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(54) Title: DISPERSING AGENTS IN NANOCOMPOSITES

Dispersing agents in nanocomposites

The present invention reletes to a composition, preferably a nanocomposite meterial, comprising (a) a synthetic polymer, especially a polycelfin, (b) a filler, especially a natural or synthetic phylicallicate or a mixture of such phylicallicates, preferably in nanoparticles, and (c) a dispereing agent prepared by controlled free radical polymerization (CFRP). The present invention relates also to a process for the prepared to 9 synthetic polymer anocomposite material which comprises met finising a mixture of a) a synthetic polymer, b) a filler, and o) a dispersing agent prepared by controlled free radical polymerization and to the use of the latter to intercelate and exfoliate a filler and disperse the filler in a synthetic polymer matrix to form a nanocomposite material.

The addition of fillers to organio materials, especially polymers, is known and is described for example in Hans Zweifel (editor), Plastics Additives Handbook, 6th Edition, pages 901-946, Hanser Publishers, Munich 2001. The use of fillers in polymers has the advantage it is possible to bring about improvement in, for example, the mechanical properties, especially the density, hardnoss, rigidity (modulus) or reduced shrinkage of the polymer.

Using extremely small filler particles (with one dimension < 200 nm), so-called nano-scale fillers, mechanical properties, heat distortion temperature stability or fiame retardant property of the polymers can be improved with a much lower concentration, typically 2 to 10 % by weight compared to 20 to 50 % by weight with the normal micro-scale filler particles. Polymers containing nano-scale fillers combine favourable mechanical properties like strength, modulus and impact, and show improved surface qualities like gloss, lower tool wear during processing and better conditions for recycling. Coatings and films comprising nano-scale fillers show improved stability, flame retardance, gas barrier properties and scratch resistance.

Nano-scale fillers possess an extremely large surface area with high surface energy. The descrivation of the surface energy and the compatibilization of the nano-scale fillers with a polymeric substrate is, therefore, even more important than with a common micro-scale filler in order to avoid aggregation during processing or conversion of the filled polymer and to reach an excellent dispersion of the nano-scaled filler in the final article.

There is a substantial recent literature on organic-inorganic nanocomposites based on clays or layered silicates such as montmorillonite and synthetic polymers, Polyolefin nanocomposites have been prepared from organic modified clays. The clays used are generally modified with alkyl or disklyl ammonium ions or amines or in a fow cases other onium ions, like for example phosphonium ions. The ammonium ion/amine additives are usually incorporated into the clay structure by a separate solution intercalation step.

These conventional organic modified clays have a number of disadvantages when used for the preparation of polyclefin nanocomposites. Ammonium salts are thermally unstable at temperatures used in polyclefin processing or may be otherwise reactive under processing conditions. These instabilities result in poor processing stability, inferior mechanical properties, discoloration, odor formation and reduced long-term stability in addition to the formation of volatile by-products.

In order to improve the polyolefin nanocomposite formation by melt processing the use of an additional compatibilizer has been proposed, most often a maleic anhydride grafted polypropylene, which in working examples is present as a major component of the final product.

M. Kewasumi et al., Macromolecules 1997, 30, 6333 – 6338 or U.S. 5,973,083 disclose that a polypropylene nanocomposite is obtained when a clay, promodified with octadeoylammonium satis, is compounded with polypropylene in the presence of polypelenin oligomers contaning polar functionality, for example maleic antiviride grafted polypropylene.

Although compatibilizers can improve the stability of nanocomposites mainly with regard to avoiding agglomeration of the filler, the other weaknesses of the nanocomposites are not improved.

It has now been found that improved synthetic polymer materials with better long term thermostability and with reduced oder and reduced undested discoloration which occurs as a result of the decomposition of the modification agents, can be propared by the use of a dispersing agent prepared by controlled free radical polymerization. The present invention therefore provides a composition comprising

- a) a synthetic polymer,
- b) a filler, and
- a dispersing agent prepared by controlled free radical polymerization (CFRP).

Preferably, component (b) is a filter in nanoparticles.

Of interest are also compositions comprising as component (b) a filler in nanoparticles or a material which produces a filler in nanoparticles under processing conditions.

Component (c) is suitable for intercalating and exfoliating a filler and dispersing the filler in a synthetic polymer matrix, especially a thermoplastic polymer, to form a nanocomposite material.

Examples of such synthetic polymers are:

1. Polymers of monocletins and dioletins, for example polypropylene, polylacbutylene, polyduta-ti-ene), polylachutylypene, polylachutylypene, polylachutylypene, polylachutylypene, polylachutylypene, polylachutylene, as well as polymers of cycloeletine, for instance of cyclopentene or norbornene, polystylene (which optionally can be crosslinked), for example high density polyethylene (HDPE), high density and high molecular weight polyethylene (HDPE-thMV), high density and ultrahigh molecular weight polyethylene (HDPE-thMV), medium density polyethylene (MDPE), linoar low density polyethylene (LLDPE), linoar low density polyethylene (LLDPE), (LLDPE) and (ULDPE).

Polyolefins, i.e. the polymers of monoolelins exemplified in the preceding paragraph, preferably polyethylene and polypropylene, can be prepared by different, and especially by the following, methods:

- a) radical polymerisation (normally under high pressure and at elevated temperature).
- catalytic polymerisation using a catalyst that normally contains one or more than one metal of groups IVb, Vb, Vib or Vill of the Periodic Table. These metals usually

have one or more than one ligand, typically oxides, halides, sicoholates, esiers, ethers, amines, alkyts, alkenyts and/or enyts that may be either rs- or -coordinated. These metal complexes may be in the free form or fixed on substrates, typically on advivated magnesium chlorids, titanhum(III) chlorids, alumina or elilion oxide. These catalysts may be soluble or insoluble in the polymerisation medium. The catalysts can be used by themselves in the polymerisation or further activators may be used, typically metal alkyts, metal hydrides, metal alkyl halides, metal alkyl oxides or metal alkylorates, said metals being elements of groups (a, its and/or III of the Periodic Table. The activators may be modified conventiently with further ester, ether, amine or silyl ether groups. These catalyst systems are usually termed Philips, Standard OII Indiana, Ziegler (-Natta), TNZ (DuPont), metallicoane or single site catalysts (SSC).

- Mixtures of the polymers mentioned under 1), for example mixtures of polypropylene with polyisobutylene, polypropylene with polyethylene (for example PP/HDPE, PP/LDPE) and mixtures of different types of polyethylene (for example LDPE/HDPE).
- 3. Copolymers of monopletins and dipletins with each other or with other vinyl monomers, for example ethylene/propylene copolymers, linear low density polyethylene (LLDPE) and mixtures thereof with low density polyethylene (LDPE), propylene/but-1-ene copolymers, propylene/isobutylene copolymers, ethylene/but-1-ene copolymers, ethylene/hexene copolymers, ethylene/methylpentene copolymers, ethylene/heptene copolymers, ethylene/octene copolymers, ethylene/vinylcyclohexane copolymers, ethylene/cycloolefin copolymers (e.g. ethylene/norbornene like COC), ethylene/1-olefins copolymers, where the 1-olefin is generated in-situ; propylene/butadiene copolymers, isobutylene/isoprene copolymers, ethylene/vinylcyclohexene copolymers, ethylene/alkyl acrylate copolymers, ethylene/alkyl methacrylate copolymers, ethylene/vinyl acetate copolymers or ethylene/acrylic acid copolymers and their salts (lonomers) as well as terpolymers of ethylene with propylene and a diene such as hexadiene, dicyclopentadiene or ethylidene-norbornene; and mixtures of such copolymers with one another and with polymers mentioned in 1) above, for example polypropylene/ethylene-propylene copolymers, LDPE/ethylene-vinyl acetate copolymers (EVA), LDPE/ethylene-acrylic acid copolymers (EAA), LLDPE/EVA, LLDPE/EAA and alternating or random polyalkylene/carbon monoxide copolymers and mixtures thereof with other polymers, for example polyamides.

4. Hydrocarbon resins (for example C_6 : C_9) including hydrogenated modifications thereof (e.g. tacktifers) and mixtures of polyalkylenes and starch.

Homopolymers and copolymers from 1.) - 4.) may have any stereostructure including syndiotectic, isotactic, heterotectic or atactic, Stereoblock polymers are also included.

- 5. Polystyrene, poly(p-methylstyrene), poly(a-methylstyrene).
- 6. Aromatic homopolymers and copolymers derived from viryl aromatic monomers including styrene, armsthylstyrene, all isomers of viryl foluene, especially p-virylloluene, all isomers of othylstyrene, propylstyrene, viryllstyhenyl, virylnaphthelane, and virylanthracone, and mixtures thereof. Homopolymers and copolymers may have any stereostructure inducting syndiotactic, isotactic, heterotactic or stactic; where stactic polymers are preferred. Stereoblock polymers are preferred.
- 6a. Copolymers Including aforementioned vinyl anomatio monomers and comonomers selected from citylene, propylene, dienes, nitriles, acids, maleti anhydrides, malelmides, vinyl aceatela and vinyl chotted or acyrilic derivatives and mixtures thereof, for example styrene/butadiene, styrene/acrylonitrile, styrene/strylone (Interpolymers), styrene/allyl methacylate, styrene/butadiene/allyl acrylate, styrene/butadiene/allyl methacylate, styrene/maletic anhydride, styrene/acrylonitrile/methyl acrylate; mixtures of hijh impact styrength of styrene copolymers and another polymer, for example a polyacrylate, a diene polymer or an citylene/propylene/diene isrpolymer; and block copolymers of styrene such as styrene/butadiene/styrene, styrene/styrene, styrene/ethylene/styrene or styrene/styrene, styrene/ethylene/styrene
- 6b. Hydrogenated aromatic polymers derived from hydrogenation of polymers mentioned under 6.), especially including polycyclohexylethylene (PCHE) prepared by hydrogenating atactic polystyrene, often referred to as polyvinylcyclohexane (PVCH).
- Hydrogenated aromatic polymers derived from hydrogenation of polymers mentioned under 6a.).

Homopolymers and copolymers may have any stereostructure including syndiotactic, isotactic, hami-isotactic or atactic; where atactic polymers are preferred. Stereoblock polymers are also included.

- 7. Graft copolymers of vinyl aromatic monomers such as styrone or c-methylstyrone, for example styrone on polybutadiene, styrene on polybutadiene-styrone or polybutadiene-styrone or polybutadiene-styrone or polybutadiene-styrone or polybutadiene; styrene and acylonitrile (or methacrytonitrile) on polybutadiene; styrene, acylonitrile and mateix anhydride or polybutadiene; styrene, acylonitrile and mateix anhydride or mateinide on polybutadiene; styrene and mateix on polybutadiene; styrene and antiplic on polybutadiene; styrene and acylonitrile on ethylene/propylene/diene terpolymers; styrene and acylonitrile on ethylene/propylene/diene terpolymers; styrene and acylonitrile on acylonitrile on polybutadiene copolymers, as well as inbutures thereof with the copolymers listed under 6), for example the copolymer mixtures known as ABS, MBS, ASS or ASS oolymers.
- 8. Halogen-containing polymers such as polychicroprene, chlorinated rubbers, chlorinated and brominated copolymer of isobutyfene-isoprene (halobutyf rubber), chlorinated or sulfa-chlorinated polyethylene, copolymers of ethylene and chlorinated ethylene, epichicro hydrin homo- and copolymers, especially polymers of halogen-containing viryl compounds, for example poly(vinyl chloride), poly(vinyl didene chloride), poly(vinyl didene chloride), poly(vinyl didene chloride), to be used as onpolymers thereof such as vinyl chloride/vinyl idene or vinylidene chloride/vinyl acetate copolymers.
- Polymers derived from α,β-unsaturated acids and derivatives thereof such as polyacrylates and polymethacrylates; polymethyl methacrylates, polyacrylamides and polyacrylonitriles, which can be impact-modified with butyl acrylate.
- 10. Copolymers of the monomers mentioned under 9) with each other or with other unsaturated monomers, for example acrylonitinel butadiene copolymers, acrylonitinel/aliky lacrylate or acrylonitinel/aliky lacrylate or acrylonitinel/aliky methacrylate/butadiene terpolymers.

- 11. Polymers derived from unsaturated alcohols and amines or the eoyl derivatives or acotals and their hydrolysis products, for example poly(viny) alcohol), poly(viny) acetalo, poly(viny) starate), poly(viny) starate), poly(viny) bolycate), poly(viny) maleate), poly(viny) bulyrat), poly(eliy) prihitale) or poly(eliy) melamino); as well as their copolymers with ofefina monitioned in 1) above.
- Homopolymers and copolymers of cyclic ethers such as poly(alkylene glycols), poly-(athylone oxide), poly(propylene oxide) or copolymers thereof with bisglycidyl ethers.
- Polyacetals such as polyoxymethylene and those polyoxymethylenes which contain othylene oxide as a comonomer; polyacetals modified with thermoplastic polyurethanes, appliets or MB9.
- 14. Polyphenylene oxides and sulfides, and mixtures of polyphenylene oxides with styrene polymers or polyamides.
- 15. Polyurethanes derived from hydroxyl-terminated polyethers, polyesters or polybutadienes on the one hand and alliphatic or aromatic polyisocyanates on the other, as well as precursors thereof.
- 16. Polyamides and copolyamides derived from diamines and dicarboxyllo acids and/or from aminocarboxyllo acids or the corresponding laclams, for example polyamide 4, polyamide 6, polyamide 66, polyamide 676, eMo, 699, eM2, 416, 12/12, polyamide 11, polyamide 12, aromatic polyamides starting from moxylene diamine and acido acid; polyamides prepared from hexamethyleneclamine and teophitalic or/and terephithaline acid and with or without an elastomer as modifier, for example poly-2-4,4,-trimethylenexamethylene terephithalined or poly-m-phonylene isophitisalamide; and also block copolymers of the aforementioned polyamides with polyetelians, clotin copolymers, ionomers or chemically bended or grafted elastomers; or with polyethers, e.g. with polyethylene glycol, polypropylene glycol or polytelramethylene glycol; as well as polyamides or copolyamides modified with EPDM or ABS; and polyamides condensed during processing (RIM polyamide systems).
- 17. Polyureas, polyimides, polyamide-imides, polyetherimids, polyesterimids, polyhydantolns and polybenzimidazoles.

- 18. Polyesters derived from dicarboxylic acids and diols and/or from hydroxycarboxylic acids or the corresponding lactones, for example polyethylene terephthalate, poly-1,4-dimethylolcyclohexane terephthalate, poly-1,4-dimethylolcyclohexane terephthalate, poly-1,6-dimethylolcyclohexane terephth
- 19. Polycarbonates and polyester carbonates.
- 20. Polyketones.
- 21. Polysulfones, polyether sulfones and polyether ketones.
- 22. Crosslinked polymers derived from aidehydes on the one hand and phenois, ureas and melamines on the other hand, such as phenoi/formaldehyde resins, urea/formaldehyde resins and melamine/formaldehyde resins.
- 23. Drying and non-drying alkyd resins.
- 24. Unsaturated polyester resins derived from copolyesters of saturated and unsaturated dicarboxylic acids with polyhydric alcohols and vinyl compounds as crosslinking agents, and also halogen-containing modifications thereof of low flammability.
- Crosslinkable acrylic resins derived from substituted acrylates, for example epoxy acrylates, urethane acrylates or polyester acrylates.
- 26. Alkyd resins, polyester resins and acrylate resins crosslinked with melamine resins, urea resins, isocyanates, isocyanates, polyisocyanates or epoxy resins.
- 27. Crosslinked epoxy resins derived from allphatic, cycloaliphatic, heterocyclic or aromatic glycidy) compounds, e.g. products of diglyddyl others of blephenol A and blephenol F, which are crosslinked with customary hardeners such as anhydrides or amines, with or without socielerators.

Blends of the aforementioned polymers (polyblends), for example PP/EPDM, Polyamide/EPDM or ABS, PVC/EVA, PVC/ABS, PVC/MBS, PC/ABS, PBTP/ABS, PC/ASA, PC/PBT, PVC/CPE, PVC/acrylates, POM/thermoplastic PUR, PC/thermoplastic PUR, PC/Macylate, POM/MBS, PPO/HIPS, PPO/PA 6.8 and copolymers, PA/HDPE, PA/PP, PA/PPO, PBTP/EC/ABS or PBT/PET/PC.

The synthetic polymers to be protected are preferably thermoplastic polymers, especially polyoletine, polystyrenes, polyamidos, polyesters, polyacrylates, most preferably polyoletins, in particular polyethylene and polypropylene or copolymers thereof with mono- and dioletins.

Preferred fillers are for example natural or synthetic phyllosilicates or a mixture of such phyllosilicates.

Of interest are compositions wherein component (b) is a natural or synthetic phylicalicate or a mkture of such phylicalicates wherein component (b) can be modified or unmodified by a modification agent such as, for example, an ammotum, an amine or a phosphonium corapound.

Fillers of special Interest are for example layered silicate claye. Of very special interest are compositions comprising as component (b) a montmorillonite, bentonite, beidellite, mica, heatorite, saponite, nontronite, sauconite, vermiculite, ledikite, magadite, kenyalte, steven-site, volkonskotile, hydrotaletice or a mixture thereof.

Component (b) can be unmodified or modified by a modification agent such as, for example, an ammonium, an amine or a phosphonium compound.

Examples of modification agents for nano-clays are for example:

Amine and ammonium compounds, for example, distearyldimethylammonium chloride, stearyldemzyldimethylammonium chloride, stearyldamine, stearyldiethoxyamine or aminododecanolo acid [commercially available as Nanofil (RTM) from Süchenle, Germany]; dimethyl ditallow ammonium, trioclylmethyl ammonium dipolyoxyethyleneallylmethyl ammonium or polyoxypropylenemethyldiethyl ammonium (commercially available as modified Somasif (RTM) from CO-OP Chemical);

octadecytamine, triethoxyeilanyl-propylamine [commercially available as Nanomer (RTM) from Nanocorj, polyalkoxylated ammonium compounds such as for example octadocyl bis[cploxyethylene(15]smine [Ethomeon (RTM) from Eastman] or octadecyl methyl bis[cploxyethylene(15]mmonium chloride [Eloquad (RTM) from Eastman] or just the corresponding free amines.

- <u>Phosphonium compounds</u>, for example tetrabutylphosphonium or octadecyl triphanyl phosphonium [continerotally available from Eastman].
- 3. <u>Others</u>, for example, triethoxy octyl silane [commercially available as Nanomer (RTM) from Nanocori, ammonium, sulfonium or pyridium compounds as disclosed for example in WO-A-01/04050 or WO-A-998/7790; block graft copolymers such as for example PEO-b-PS or poly-4-vinylpyridine-b-PS; or solvents for swelling such as for example "butyrolactions, 2-pyrrolidons, dimethylsulfoxide, diglyme, tetrahydrofuran or furfuryl alcohol.

Compositions which are of interest include those comprising as component (o) a random, block, star or comb copolymer prepared by controlled free radical polymerization.

Such dispersing agents prepared by controlled free radical polymerization are known. Their preparation is disclosed for example in C. Auschra et al., Progress in Organic Coalings 45, 83 – 93 (2002).

The preparetton of the dispersing agents by controlled free radical polymerization (CFRP) can preferally occur by four sulfable routes:

- Polymerization in the presence of alkoxyamine initiator/regulator compounds;
- Polymerization in the presence of a stable nitroxyl free radical and a radical initiator (source of free radicale);
- a3) Polymerization under atom transfer radical polymerization (ATRP); or
- a4) RAFT polymerization which refers to a method of polymer synthesis by radical polymerization using chain transfer agents which react by reversible addition fragmentation chain transfer.

For example U.S. 4,581,429 or EP-A-0 621 878 discloses the preparation of block copolymers by method a1).

For example EP-A-0 621 878 discloses the preparation of block copolymers by method a2).

For example WO-A-01/51534 discloses the preparation of comb copolymers by the ATRP method e3).

For example WO-A-99/31144 discloses the preparation of block copolymers by method a4).

Preferred chain transfer agents include thiocarbonylthic chain transfer agents of formulae I, II and IIII, and macromonomers of formula IV and a combination thereof,

$$(1) \qquad s = c \begin{cases} s - R \\ z \end{cases} \qquad \left[s = c \begin{cases} s - \frac{1}{2} - R^{t} \end{cases} \right]_{n}$$

wherein R, R¹ and P ere free radical leaving groups; and wherein,

R is selected from the group consisting of unsubstituted or substituted Cr-Crealityi, unsubstituted or substituted Cr-Crealityi, unsubstituted or substituted carbocycle ring, unsubstituted or substituted carbocycle ring, unsubstituted or substituted unsaturated carbocycle ring, unsubstituted or substituted unsaturated carbocycle ring, unsubstituted aromatic carbocycle ring, unsubstituted or substituted unsaturated heterocycle ring, unsubstituted or substituted unsaturated heterocycle ring, unsubstituted or substituted or subs

Z is selected from the group consisting of hydrogen, chlorine, unsubstituted or substituted C_{1} - C_{1} -galley), unsubstituted or substituted anyl, unsubstituted or substituted C_{2} - C_{1} - C_{1} - C_{2} - C_{3} - C_{4} - C_{4} - C_{4} - C_{5} - C_{4} - C_{5} -

heterocyclic ring, unsubstituted or substituted Cr-Creatively, unsubstituted or substituted arythio, unsubstituted or substituted Cr-Creatively, unsubstituted or substituted are substituted or substituted are substituted or substit

R1 is an n-valent molety derived from R;

Z' is an m-valent molety derived from a species selected from the group consisting of unsubstituted or substituted cr.O₁₆ allyl, unsubstituted or substituted anyl and a third polymer chain having a number evenega degree of polymerization in the range of 5 to 1000 wherein the connecting moleties in said polymer chain are selected from the group consisting of Cr-O₁₆-allphatic carbon, aromatic carbon, oxygen, sulfur and a combination thereof.

P is an oilgomer or polymer having a number average degree of polymerization in the range of 2 to 1000 or said first polymer chain;

X is selected from the group consisting of unsubstituted or substituted aryl, C₁-C₁₁-alkoxycarbonyl, unsubstituted or substituted aryloxycarboyl, carboxy, unsubstituted or substituted aryloxycarbonyl, C₁ to C₁₅ carboxy, and cyano;

n and m are integers in the range of 1 to 6; and

wherein the substituents for the substituted groups for R, R¹, Z, z¹ and X are independently selected from the group consisting of hydroxy, teritary amino, halogan, cyano, epoxy, carboxyflo acid, alkoxy, alkyl having 1 to 32 carbon atoms, alkynyl having form 2 to 32 carbon atoms, esturated carbocyclic rings having 3 to 14 carbon atoms, unsaturated carbocyclic rings having 4 to 14 carbon atoms, aromatic carbocyclic rings having 6 to 14 carbon atoms, saturated heterocyclic rings having 3 to 14 carbon atoms, unsaturated heterocyclic rings having 3 to 14 carbon atoms and aromatic carbocyclic rings having 6 to 14 carbon atoms.

By a "polymer chain" referred to above for R, R¹, Z, Z¹ and P is meant conventional condensation polymers, such as polyesters [for example, polycaprolacione, poly(elhytene tetephrithate), poly(actio acid)), polycarbonates, poly(alkytene oxide)s [for example, poly(elhytene oxide), poly(tetramethytene oxide)), in/tone, polyuretitanes or chain polymers such as those formed by coordination polymerization (for example polyethylene, polypropylene), radical polymerization (for example, polymeth)socylates and polyetyrenics, anionto polymerization (for example, polystyrene, polybutadiene), cationia polymerization (for example, polylsobutylene) and group transfer polymerization (for example, poly(meth)acrylates).

The groups R, R¹, Z and Z¹ are preferably chosen so as to be appropriate for use with the desired monomers or monomer combination and the polymerization method as disclosed in WO-A-98/01478, WO-A-98/05099 and WO-A-98/31144, all of which are incorporated herein by reference.

The synthesis of block copolymers by RAFT polymerization can be carried out in emulsion, solution or suspension in either a batch, semi-batch, continuous, or feed mode. For lowest polydispersity block copolymers, the RAFT sgent is typically added before polymerization is commenced. For example, when carried out in a batch mode in solution, the reactor is typically charged with a RAFT agent and a monomer, or a polymerization medium plus the monomer. Block copolymers with broader, yet controlled, polydispersity or with multimodal molecular weight distribution can be produced by controlled addition of the RAFT agent over the course of the polymerization process.

In the case of emulsion or suspension polymerization, the polymerization medium will often be predominantly water and the conventional stabilizers, dispersants and other additives can be present.

For solution polymerization, the polymerization medium can be chosen from a wide range of media to suit the monomer(s) being used. For example, aromatic hydrocarbons, such as, petroleum naphtha or xylenes; ketones, such as, methyl amyl ketone, methyl isobutyl ketone, methyl ketone or acelone; esters, such as, butyl acelate or hexyl acelate; and glycol ether esters, such as, propylene glycol monomethyl ether acetate.

The use of feed polymerization conditions allows the use of RAFT agents with lower chain transfer constants and allows the synthesis of block copolymers that are not readily achieved using batch polymerization processes. If the RAFT polymerization is carried out as a feed system, the reaction can be carried out as follows:

The reactor is charged with the chosen polymerization medium, the RAFT agent and optionally a portion of the monomer mixture. Into a separate vessel is placed the remelting monomer mixture. The free radical initiator is dissolved or suspended in polymerization medium in another separate vessel. The medium in the reactor is heated and stirred while the monomer mixture plus the medium and the hititator plus the medium, is introduced, for example by a syringe pump or other pumping device. The rate and duration of feed is largely determined by the quantity of the solution, the desired monomer/RAFT agent/initiator ratio and the rate of the polymerization. When the feed is complete, heating may be continued for an additional period.

Following completion of the polymerization, the block copolymer can be isolated by stripping off the medium and unreacted monomer(s) or by predipitation with a non-solvent. Alternatively, the polymer solution/emulsion can be used as such, if appropriate to its application.

Block and multi-block and gradient copolymers may be prepared by varying the rate of monomer(s) addition and/or by varying the sequence in which the monomer(s) may be added to the polymerization medium. Gradient block may also be prepared in a one-slap process by making use of the inherent difference in reactivity of the monomer(s). For gradient block copolymers, it is often desirable to pick comonomers with disparate reactivity ratios, For example, mailed anhydride and syrone or (meth)acrylates.

By appropriate selection of the groups R, R¹, Z, Z¹, P and X of the RAFT chain transfer agent of formulae I-IV above, block copolymers with specific and functionalities can be readily produced, Block copolymer may be prepared by starting with an existent polymer chain prepared by means other than RAFT polymerization and then modifying the end or other functionality such that the polymer chain is incorporated in a compound of formulae I-IV so as to create a macromolecular RAFT agent that can be chain extended by RAFT oplymerization.

The "source of free radicals" or "initiator" suitable for use in the present invention includes those compounds that provide initiating radicals that add to monomers to produce propagating radicals.

The amount of initiator used depends upon the desired polydispersity, molecular weight and polymer structure of the resulting polymer. However, generally less than 10 weight percent based on the total amount of monomer mixture is used. Preferably the amount of initiator used is in the rance of 0,001 to 5.0 weight percent.

The source of initiating radicals includes such sources as the thermally induced homolytic scission of a suitable initiator, such as percoides, percoyasters, or azo compounds; the spentaneous generation from monomer, such as elsowers initiating systems or high energy radiation source, such as electron beam, X-or y-radiation or a combination of these methods. The initiating system is chosen such that under the reaction conditions there is substantially no adverse interaction of the initiator or the initiating radicals with the RAFT agent under the reaction conditions. The initiator should also have the regulatic solubility in the reaction medium or the monomer mixture.

Examples of suitable sources of free radicals for the process include azo compounds and peroxides such as for example 2,2'-azobis(isobutyronitrile), 2,2'-azobis(2-oyano-2-butane), dimethyl 2,2'-azobis(methyl isobutyrate), 4,4'-azobis(4-cyanopentanoic acid), 4,4'-azobis(4cyanopentan-1-ol), 1,1'-azobis(cyclohexanecarboniirile), 2-(t-butylazo)-2-cyanopropane, 2,2'-azobis[2-methyl-N-(1,1)-bis(hydoxymethyl)-2-hydroxyethyl] propionamide, 2,2'-azobis[2methyl-N-hydroxyethyl)]-propionamide, 2,2'-azobis(N,N'-dimethylenelsobutyramidine) dihydrochloride, 2,2'-azobis(2-amidinopropane) dihydrochloride, 2,2'-azobis(N,N'-dimethyleneisobutyramine), 2,2'-azobis(2-methyl-N-[1,1-bis(hydroxymethyl)-2-hydroxyethyl] propionamide), 2,2'-azobis(2-methyl-N-[1,1-bis(hydroxymethyl) ethyl] propionamide), 2,2'-azobis[2methyl-N-(2-hydroxyethyl) propionamide], 2,2'-azobls(Isobutyramide) dihydrate, 2,2'-azobls-(2,2,4-trimethylpentane), 2,2'-azobis(2-methylpropane), tert-butyl peroxyacetate, tert-butyl peroxybenzoate, t-butyl peroxyoctoate, tert-butyl peroxyneodecanoate, tert-butylperoxy isobulyrate, tert-amyl peroxypivalate, tert-butyl peroxypivalate, di-isopropyl peroxydicarbonate, dicyclohexyl peroxydicarbonate, dicumyl peroxide, dibenzoyl peroxide, dilauroyl peroxide, potassium peroxydisulfate, ammonium peroxydisulfate, di-t-butyl hyponiirite, or dicumyl hyponitrite.

Photochemical Initiator systems are chosen to have the requisite solubility in the reaction medium or monomer mixture and have an appropriate quantum yield for radical production under the conditions of the polymorization. Examples include benzon derivatives,

benzophenone, acyl phosphine oxides, and photo-redox systems. Free radicals may also be derived by direct photolysis of the compound of formula I-III.

Redox initiator systems are chosen to have the requisite solubility in the reaction medium or monomer mixture and have an appropriate rate of radicel production under the conditions of the polymorization; these initiating systems can include combinations of the following oxidants and reductants: Oxidants: potassium peroxydisulfate, hydrogen peroxide, and tertbulyt hydroperoxide. Reductants: Iron (II), titanium (III), potassium thiosulfito, and potassium bisulfite.

Other suitable Initiating systems are described in recent texts. See, for example, Moad and Solomon "The Chemistry of Free Radical Polymerization", Pergamon, London, 1995, pp 53-

Compositions which are of interest include those comprising as component (a) a block or comb copolymer having at least one clay compatible block and at least one matrix polymer compatible block prepared by controlled free radical polymertzation.

Preferably, a clay competible block is based on hydrophilic or "clayophilic" monomers which contain polar groups such as for example ether [-0-], amide [$\stackrel{Q}{-c}$ -\lambda], thioamide

 $[-\overset{\circ}{c}-\overset{\circ}{v}] \quad \text{I. nitrile, anthydride, hydroxy, amine, pyridine, ammonium and phosphonium in appropriate proximity.}$

Preferred hydrophilio or "clayophilio" monomers are for example PEO acrylate, 1-vinyl-2-pyrrolldinone, N,N-dimethyl acrylamide, acrylonlitrile, maleic anhydride, hydroxyethyl acrylate, hydroxyropyl acrylate, tert-butyl c-hydroxymethacrylate, N,N-dimethylaminoethyl acrylate, 4-vinylbenzyldihydroxyethylamine, 4-vinylpyddino or 4-vinylbenzyltributylphosphonium chlorida. The molecular weight of the acrylic copolymer is preferably between 1000 and 100000 and the polydispersity between 1.05 and 3.0. More preferably the molecular weight is between 2000 and 20000 and the polydispersity is between 1.05 and 2.0.

Copolymers of the above polymers with other monomers may also be used.

The polar (hydrophillo) or "clayophillo" segments may also be derived from a polar condensation or other polymers such as for oxample poly(ethylene oxide).

Preference is given therefore to compositions comprising as component (c) a block or comb acrylic copolymer prepared by controlled free radical polymerization.

The block or comb copolymers prepared by controlled free radical polymerisation can also be described as polymeric dispersing or solvating agents having amphiphilic properties. They have polar and nonpolar groups in the same molecule and they are, for example, dispersing or solvating agents based on polyethylene glycols (PEG), polyacrylates, polyeting agents based on polyethylene glycols (PEG), polyacrylates, polyeting agents based on polyethylene glycols (PEG), polyacrylates, polyeting agents based on polyethylene containing at least one block copolymer based on scrylate, acrylic acid or methacrylate.

Block copolymers are, for example, diblock copolymers (A-B type) or triblock copolymers (A-B-A or A-B-C type) and so-called tapered structures.

Diblock copolymers (A-B type) are, for example, poly(etyrene-b-methylmethacrylate), poly(etyrene-b-tert-butylmethacrylate), poly(butadiene-b-tert-butylmethacrylate), poly(butylmethacrylate), poly(butylmetha

methyl methacrylate), poly(tert-butyl methacrylate-b-tert-butyl acrylate), poly(tert-butyl methacrylate-b-2-vinylpyridine), poly(tert-butyl methacrylate-b-2-vinylpyridine), poly(tert-butyl acrylate-b-2-vinylpyridine), poly(tertylane-b-methyl methacrylate), poly(tetylane-b-2-vinylpyridine) or poly(tetylane-b-4-vinylpyridine).

Triblock copolymers of the A-B-A type are, for example, poly(methyl methacrylate-b-styreneb-methyl methacrylate), poly(tert-bulyl methacrylate-b-styrene-b-tert-butyl methacrylate), poly(tert-butylacrylate-b-styrene-b-tert-butylacrylate), poly(2-vinylpyridine-b-styrene-b-tertbulyl acrylate), poly(4-vinylpyridine-b-styrene-b-4-vinylpyridine), poly(butadlene[1,2addition]-b-styrene-b-butadiene[1,2-addition]), poly(butadiene[1,4-addition]-b-styrene-bbutadlene[1,4-addition]), poly(styrene-b-butadlene[1,4- and 1,2-addition]-b-styrene), poly(methyl methacrylate-b-butadiene[1,4- or 1,2-addition]-b-methyl methacrylate), poly(tertbutyl methacrylate-b-methyl methacrylate-b-tert-butyl methacrylate), poly(tert-butyl acrylateb-methyl methacrylate-b-tert-butyl acrylate), poly(methyl methacrylate-b-2-vinylpyridine-bmethyi methacrylate), poly(4-vinylpyridine-b-methyl methacrylate-b-4-vinylpyridine), poly(methyl methacrylate-b-tert-butyl acrylate-b-methyl methacrylate), poly(methyl methacrylate-b-n-butyl acrylate-b-methyl methacrylate), poly(tert-butyl methacrylate-b-tertbutyl acrylate-b-lert-butyl melhacrylate), poly(2-vinylpyridine-b-tert-butyl acrylate-b-2vinylpyridine), poly(4-vinylpyridine-b-tert-butyl acrylate-b-4-vinylpyridine), poly(styrene-b-nbutyl acrylate-b-styrene), poly(styrene-b-ethyl acrylate-b-styrene), poly(styrene-b-ethylene-bstyrene), poly(styrene-b-butylene-b-styrene), poly(ethylene oxide-b-styrene-b-ethylene oxide), poly(styrene-b-ethylene oxide-b-styrene) or poly(styrene-b-acrylic acid-b-styrene).

Triblock copolymers of the A-B-C type are, for example, poly(styrene-b-butadiane-b-2-viny) pyridina), poly(styrene-b-butadiane-b-4-winy)pyridina), poly(styrene-b-butadiane-b-4-winy)pyridina), poly(styrene-b-a-viny)pyridina), poly(styrene-b-2-viny)pyridina-b-d-4-winy)pyridina), poly(styrene-b-a-winy)pyridina-b-d-winy)

Particularly sultable dispersing or solvating agents having amphiphilic properties are, for example, poly(butadiene-b-methyl methacrylate), poly(isoprene-b-methyl methacrylate), poly(isoprene-b-methyl methacrylate), poly

(ethylene-b-methyl methacrylate), poly(styrene-b-4-vinylpyridine), poly(styrene-2-b-vinylpyridine), dine), poly(styrene-b-n-butyl acrylate), poly(styrene-b-tert-butyl acrylate), poly(styrene-bsodium acrylate), poly(styrene-b-acrylic acid), poly(methyl methacrylate-b-sodium acrylate), poly(methyl methacrylate-b-sodium methacrylate), poly(ethylene oxide -b-s-caprolactone), poly(2-vinylpyridine-b-ethylene oxide), poly(butadlene-b-ethylene oxide), poly(butadlene-bsodium acrylate), poly(ethylene-b-ethylene oxide), poly(ethylene-b-propylene oxide), poly-(styrene-b-ethylacrylate-b-styrene), poly(ethylene oxide-b-styrene-b-ethylene oxide), poly-(styrene-b-acrylic acid-styrene), poly(styrene-b-butadiene-b-methyl methacrylate), poly(styrene-b-acrylic acid-styrene), rene-b-vinylpyridine-b-ethylene oxide), poly(styrene-b-4-vinylbenzolo acid), poly(styrene-bpolyglyoldyl-methacrylate), poly(ethylene-b-glyoldyl methacrylate), poly(propylene-b-acryllo acid), poly(ethylene-b-acrylic acid), poly(propylene-b-maleic anhydride), poly(ethylene-bmaleic anhydride), poly(slyrene-b-maleic anhydride), poly(methacrylic acid)-poly(alkylene oxide) block copolymers, for example according to EP-A-0 859 028, polysiloxane-polyoxyalkylene, copolymers of maleates and styrene or styrene derivatives, for example according to EP-A-0 791 024, polystyrene-polysiloxane block copolymers, polyacrylate-polysiloxene block copolymers and cyclosiloxane-radiale copolymers, prepared e.g. using ATRP technology according to EP-A-0 870 774, methylacrylate-styrene copolymer, methylmethacrylatestyrene, polybutadiene-methylacrylates, prepared by nitroxyl-initiated radical polymerisation according to EP-A-0 135 280.

The polyalkylene oxides are preferably polyethylene oxide, polypropylene oxide and polyhutviene oxide.

Sultabla block copolymers are, for example, polyacrylate/polystyrene, polymethacrylate/polysthylene oxide, polyacrylate/polystyrene, polyurinyl acetale/ polyethylene polyacrylate/polybutadiene, polyacrylate/polybutadiene, polyacrynethacrylate, polyethylene/polymethylmethacrylate, polyethylene/polyethylene/polyethylene/polyethylene/polyethylene/polyethylene/polyethylene/polyethylene/polyethylene/polyethylene/polyethylene/polyethylene/polyethylene/polye

Particularly autitable disporaing or solvating agents having amphiphilic properties are, for example, poly(styrene-bis-sodium medylate), poly(styrene-bis-sodium methacytate), poly(styrene-bis-dium methacytate), poly(styrene-bis-di-methyl-d-vin/pipridinium lodde), poly(soprene-bis-div), methyl-d-vin/pipridinium lodde), poly(soprene-bis-dhylene oxide), poly(methylmethacry-methyl-d-vin/pipridinium lodde), poly(soprene-bis-dhylene oxide), poly(methylmethacry-methyl-d-vin/pipridinium).

late-bis-sodium acytate), poly(methylmethacrytate-bis-sodium methacrytate), poly(methylmethacrytate-bis-ethylene oxide), poly(tert-butylmethacrytate-bis-ethylene oxide), poly(methylmethacrytate-bis-bis-bis-datato), poly(2-vtnylypridina-bis-ethylene oxide), poly(2-vtnylypridina-bis-ethylene oxide), poly(butadiene-bis-eodium exheroriate), poly(butadiene-bis-bis-bis-bis-bis-bis-bis-ethylene oxide), poly(butadiene-bis-bis-ethylene oxide), poly(butadiene-bis-bis-ethylene oxide) or poly(ethylene-bis-propylene oxide).

Other likewise preferred dispersing or solvating agents based on polyacrylates are described, inter alia, in U.S. 5,133,899.

Preferably, component (b) is present in the composition in an amount of from 0.1 to 40 %, more preferably 0.5 to 20 %, most preferably from 1 to 10 %, based on the weight of the synthetic polymer [component (a)].

Preferably, component (c) is present in the composition in an amount of from 0.1 to 20 %, more preferably 0.1 to 15 %, most preferably from 0.1 to 10 %, based on the weight of the synthetic polymer [component (a)].

In addition to components (a), (b) and (c) the novel compositions may comprise further additives, such as, for example, the following:

1. Antioxidants

1.1. Alkvleted monophenols, for exemple 2,6-di-tert-butyl-4-methylphenol, 2-butyl-4,8-dimothylphenol, 2,6-di-tert-butyl-4-dihylphenol, 2,6-di-tert-butyl-4-nebutylphenol, 2,6-di-tertbutyl-4-la-butylphenol, 2,6-di-cydopentyl-4-methylphenol, 2-(a-methylphenol, 2,6-di-tertbutyl-4-methylphenol, 2,6-di-tert-butyl-4-methylphenol, 2,6-di-tert-butyl-4methylphenol, 2,6-di-tert-butyl-4methylphenol, 3,6-di-tert-butyl-4methylphenol, 3,6-di-tert-butyl-4methylphenol, 3,6-di-tert-butyl-4methylphenol, 2,4-dimethyl-6-(1'-methyltindec-1'-yl)-phenol, 2,4-dimethyl-6-(1'methylpheladec-1'-yl)-phenol, 2,4-dimethyl-6-(1'-methyltindec-1'-yl)-phenol and mbdures
methylphenol, 2,4-dimethyl-6-(1'-methyltindec-1'-yl)-phenol and mbdures

- Alkytthiomethylchenols, for example 2.4-dioctylthiomethyl-6-tert-butylphenol, 2.4-dioctylthiomethyl-6-methylphenol, 2.4-dioctylthiomethyl-6-methylphenol, 2.6-diodecylthiomethyl-4-nonvlohenol.
- 1.3. Hydrogulinonea and alikylated hydrogulinonea, for axample 2,6-di-tert-butyl-4-methoxy-phenol, 2,6-di-tert-butyl-hydrogulinone, 2,5-di-tert-amylinydrogulinone, 2,6-di-tert-butyl-4-hydroxyanisole, 3,5-di-tert-butyl-4-hydroxyanisole, 3,5-di-tert-butyl-4-hydroxyanisole, 3,5-di-tert-butyl-4-hydroxyanisole, 3,5-di-tert-butyl-4-hydroxyanisole, 3,5-di-tert-butyl-4-hydroxyanisole, 3,5-di-tert-butyl-4-hydroxyanisole
- 1.4. Tocopherols, for example ω -tocopherol, β -tocopherol, γ -tocopherol, δ -tocopherol and mixtures thereof (Vitamin E).
- 1.6. Hvdroxvlated thiodiphenvi ethers, for example 2,2-thiobis(6-tert-butyl-4-methylphenol), 2,2-thiobis(4-octylphenol), 4,4-thiobis(6-tert-butyl-3-methylphenol), 4,4-thiobis(6-tert-butyl-2-methylphenol), 4,4-thiobis(3,6-di-soc-amylphenol), 4,4-bis(2,6-dimethyl-4-hydroxyphenyl) disulfide.

meltryiphenyi)dioyclopentadiene, bis[2-(3'-tert-butyl-2'-hydroxy-5'-meltryibenzyi)-6-tert-butyl-4-mettryiphenyij terephthelate, 1,1-bis[3,5-dimethyl-2-hydroxyphenyi)butane, 2,2-bis[3-dimethyl-2-hydroxyphenyi)propane, 2,2-bis[6-tert-butyl-4-hydroxy-2-meltryiphenyi)-4-meltryiphenyi-4-melt

- 1.7. O. N. and S.-benzyl compounds, for example 3,5,3',5'-letra-tert-bufy4.4'-dihydroxy-dibenzyl ether, octadecyl-4-hydroxy-3,5-dimethylbenzyl mercaploacelate, tridecyl-4-hydroxy-3,5-di-dr-bufyl-benzyl mercaploacelate, tris(3,5-di-tert-bufy4-hydroxy-benzyl)amine, bis(4-tr-bufy4-hydroxy-2,6-dimethylbenzyl) dilhoterephthalate, bis(3,5-di-tert-bufy4-hydroxy-benzyl) suffice, isocofyl-3,5-di-tert-bufy4-hydroxy-benzyl mercaploacelate.
- 1.8. <u>Hydroxybenzylated malonates</u>, for example dioctadeoyl 2,2-bis(3,5-di-lart-butyl-2-hydroxybenzyl)malonate, dioctadeoyl 2(3-tert-butyl-4-hydroxy5-mathylbenzyl)malonate, didodecylmercaptoethyl 2,2-bis(3,5-di-lart-butyl-4-hydroxybenzyl)malonate, dt-[4-(1,1,3,3-tertamethylbutyl)phenyl] 2,2-bis(3,5-di-lart-butyl-4-hydroxybenzyl)malonate.
- 1.9. Hydroxybepzyl eromatic compounds, for example 1,3,5-tris(3,5-di-tert-butyl-4-hydroxybenzyl)-2,4,6-trimethylbenzene, 1,4-bis(3,5-di-tert-butyl-4-hydroxybenzyl)-2,3,5,6-tetramethylbenzene, 2,4,6-tris(3,5-di-tert-butyl-4-hydroxybenzyl)phenol.
- 1.10. Triazine compounds, for example 2,4-bisocty/imercapto-8-(3,5-di-tert-butyl-4-hydroxy-anllino)-1,3,5-triazine, 2-octy/imercapto-4,6-bis(3,5-di-tert-butyl-4-hydroxy-anllino)-1,3,5-triazine, 2-octy/imercapto-4,6-bis(3,5-di-tert-butyl-4-hydroxy-phenoxy)-1,3,5-triazine, 2,4,6-tris-(3,5-di-tert-butyl-4-hydroxy-phenoxy)-1,2,3-triazine, 1,3,5-tris(3,5-di-tert-butyl-4-hydroxy-phenoxy-2,6-dimethylbenzyl) isocyanurate, 2,4,6-tris(3,5-di-tert-butyl-4-hydroxy-phenylethyl)-1,3,5-triazine, 1,3,5-tris(3,5-di-tert-butyl-4-hydroxy-phenylethyl)-1,3,5-triazine, 1,3,5-tris(3,5-di-tert-butyl-4-hydroxy-phenylethyl)-1,3,5-tris(3,5-di-tert-butyl-4-hydroxy-phenylethyl-1,3,5-tris(3,5-di-tert-butyl-4-hydroxy-phenylethyl-1,3,5-tris(3,5-di-tert-butyl-4-hydroxy-phenylethyl-1,3,5-tris(3,5-di-tert-butyl-4-hydroxy-phenylethyl-1,3,5-tris(3,5-di-tert-butyl-4-hydroxy-phenylethyl-1,3,5-tris(3,5-di-tert-butyl-4-hydroxy-phenylethyl-1,3,5-tris(3,5-di-tert-butyl-4-hydroxy-phenyl-1,3,5-tris(3,5-di-tert-butyl-4
- 1.11. Benzylphosohonates, for example dimethyl 2,5-dl-tert-butyl-4-hydroxybenzyl-phosphonate, dleithyl 3,5-dl-tert-butyl-4-hydroxybenzylphosphonate, dloctadecyl 3,5-dl-tert-butyl-4-hydroxybenzylphosphonate, dloctadecyl 5-fert-butyl-4-hydroxy-3-methylbenzyl-phosphonate, calcium sail of 3,5-dl-tert-butyl-4-hydroxybenzyl-phosphonate celd monoethyl saler.
- 1.12. Acylaminophenols, for example 4-hydroxylaurio acid anilide, 4-hydroxystearic acid anilide, N-(3,5-di-lert-butyl-4-hydroxyphenyl)carbamic acid octyl ester.

1.13. Esters of B-(3.5-di-lett-buty4-hydroxyphenyl)proptonio acid with mone- or poly-hydric alcohols, for example with methanol, otheroit, n-octanol, isoocianol, citadecanol, 1,6-hoxanediol, 1,9-nonanediol, ethyleno glycol, 1,2-propanediol; n-oopentyl glycol, thiodiethyleno glycol, diethyleno glycol, tethyleno glycol, pentaerythritol, tris(hydroxyethyl) isooyanurate, N,N*-bis(hydroxyethyl)oxalo acid dlamide, 3-thiaundecanol, 3-thiapentadecanol, trimethylthexanediol, trimethylolpropane, 4-hydroxymethyl-1-phospha-1,6-f-droxabbyologi-2.2 glociano.

1.14. Esters of B-(6-tert-buty-4-hydroxy-3-methylohenyl)proplants acid with mano- or polyhydrid alcohols, for example with methanol, ethanol, n-ctanol, isocatanol, o-daddecanol, 1,6-hoxanediol, 1,9-nonanediol, ethylene glycol, 1,2-propanediol, neopeniyl glycol, intellylene glycol, prehipylene glycol, prehipylene, acid diamide, 3-thisundecanol, 3-thisundecanol, itimathylhoxanediol, trinethylolpropano, 4-hydroxymethyl-1-phosphatelyl-1-glycolyl-1,1-dimethylethyl-1-glycolyl-1,4-dimethylethyl-1-glycolyl-1,4-dimethyl-1-glycolyl-1,4-dimethyl-1-glycolyl-1,4-dimethyl-1-glycolyl-1,4-dimethyl-1-glycolyl-1,4-dimethyl-1-glycolyl-1,5-glyc

1.16. Esters of 6.43.6-diovolohexvi-4-hydroxynshenviloropionio acid with mono- or poly-hydrio alcohols, for example with methanol, ethanol, octanol, octadecanol, 1.6-hexanediol, 1,9-nonanediol, ethylene glycol, 1,2-propanediol, neopenyly glycol, thiodiethylene glycol, delthylene glycol, pentaerythritol, tris(hydroxyethyl) isocyanurala, N,N-bis(hydroxyethyl)oxalic acid diamide, 3-hisundecanol, 3-thiapentadocanol, trimethylioxanediol, trimethyliopropane, 4-hydroxymethyl-1-phospha-2.6,7-trioxabloyclo(2.2.2)cotano.

1.18. Esters of 3.8-dl-tert-butyl-4-hydroxyohonylacetto acid with mono- or poly-hydro alcohols, for example with methanol, ethanol, octand, octando, odadecanol, 1,6-hexandelol, 1,9-nonandelol, ethylens glycol, 1,2-propanedlol, neopentyl glycol, thiodiethylens glycol, idethylens glycol, pentaerythritid, tisthydroxyethyl) isocyanurate, N,N-bis(hydroxyethyl)oxalic acid diamide, 3-thiaundecanol, 3-thiapentadecanol, trimethyletropane, 4-hydroxymethyl-1-phospha-2,6,7-trioxabley/old2,2,2lodane.

1.17. Amides of B-(3.5-di-tert-butyl-4-hydroxyphenyllproplonic acid, for example N,N-bis(3.5-di-tert-butyl-4-hydroxyphenylproplonyl)hexamethylenedlamide, N,N-bis(3.5-di-tert-butyl-4-hydroxyphenylproplonyl)hydrazide), N,N-bis(2.5-di-tert-butyl-4-hydroxyphenylproplonyl)hydrazide), N,N-bis(2.5-di-tert-butyl-4-hydroxyphenylproplonyl)hydrazide), N,N-bis(2.6-di-tert-butyl-4-hydroxyphenylproplonyloxybityl)kymemide (Naugard*N,L-1 from Unfroyal).

1.18. Ascorbic acid (Vitamin C).

1.19. Amine-type antioxidants, for example N,N'-di-isopropyi-p-phenylenediamine, N,N'-disec-butyl-p-phenylenediamine, N,N'-bis(1,4-dimethylpentyl)-p-phenylenediamine, N,N'-bis(1ethyl-3-methylpentyl)-p-phenylenediamine, N,N'-bis(1-methylheptyl)-p-phenylenediamine, N,N'-dicyclohexyl-p-phenylenediamine, N,N'-diphenyl-p-phenylenediamine, N,N'-di(2naphthyl)-p-phenylenediamine, N-Isopropyi-N-phenyl-p-phenylenediamine, N-(1,3-dimethylbutyl)-N'-phenyl-p-phenylenediamine, N-(1-methylheptyl)-N'-phenyl-p-phenylenediamine, Ncyclohexyl-N-phenyl-p-phenylenediamine, 4-(p-toluenesulfonamido)-diphenylamine, N,N'dimethyl-N.N'-di-sec-butyl-p-phenylenediamine, diphenylamine, N-altyldiphenylamine, 4-isopropoxydiphenylamine, N-phenyl-1-naphthylamine, N-(4-tert-octylphenyl)-1-naphthylamine, N-phenyl-2-naphthylamine, octylated diphenylamine, for example p,p'-di-tert-octyldiphenylamine, 4-n-butylaminophenol, 4-butyrylaminophenol, 4-nonanoylaminophenol, 4-dodecanoylaminophenol, 4-octadecanoylaminophenol, di(4-methoxyphenyl)amine, 2,6-ditert-butyi-4-dimethylaminomethylphenol, 2,4'-diaminodiphenylmethane, 4,4'-diaminodiphenylmethane, N,N,N',N'-tetramethyl-4,4'-diaminodiphenylmethane, 1,2-di[(2-methylphenyl)amino]ethane, 1,2-di(phenylamino)propane, (o-tolyl)-biguanide, di[4-(1',8'-dimethylbutyl)phenyl[amine, tert-octylated N-phenyl-1-naphthylamine, mixture of mono- and dialkylated tert-butyl-/tert-octyl-diphenylamines, mixture of mono- and di-alkylated nonyldiphenylamines, mixture of mono- and di-alkylated dodecyldiphenylamines, mixture of mono- and di-alkylated isopropyl-/isohexyl-diphenylamines, mixtures of mono- and di-2.3-dihydro-3.3-dimethyl-4H-1,4-benzothlazine, tert-butyldiphenylamines. phenothlazine, mixture of mono- and di-alkylated tert-butyl-/tert-octyl-phenothlazines, mixture of mono- and di-alkylated tert-octylphenothlazines, N-allylphenothlazine or N,N,N',N'-tetraphenyl-1,4-diaminobut-2-ene.

2. UV absorbers and light stabilizers

2.1. 2-(2'-Hydroxyphenyl)-benzotriazoles, for example 2-(2'-hydroxy-5'-methylphenyl)-benzotriazole, 2-(3',5'-di-tert-butyl-2'-hydroxyphenyl)-benzotriazole, 2-(6'-tert-butyl-2'-hydroxyphenyl)-benzotriazole, 2-(2'-hydroxy-5'-(1,1,3,3-tetramethylbutyl)-phenyl)-benzotriazole, 2-(3',5'-2-(3'-tert-butyl-2'-hydroxy-5'di-tert-butyl-2'-hydroxyphenyt)-5-chlorobenzotriazole, 2-(3'-sec-bulyi-5'-tert-bulyi-2'-hydroxyphenyi)methylphenyl)-5-chlorobenzotriazole, 2-(3',5'-dl-tert-amyl-2'-2-(2'-hydroxy-4'-octyloxyphenyl)-benzolrlazole, benzotriazole, 2-(3',5'-bls(α,α-dimethylbenzyl)-2'-hydroxyphenyl)-benzohydroxyphenyl)-benzotriazole, triazole, 2-(3'-tert-butyl-2'-hydroxy-5'-(2-octyloxycarbonylethyl)phenyl)-5-chlorobenzotriazole, 2-(3'-tert-butyl-5'-[2-(2-ethylhexyloxy)carbonylethyl]-2'-hydroxyphenyl)-5-chlorobenzotriazola, 2-(3'-tert-butyl-2'-hydroxy-5'-(2-methoxycarbonylethyl)phenyl)-5-chlorobenzotriazole, 2-(3'tert-butyl-2'-hydroxy-5'-(2-methoxycarbonylethyl)phenyl)-benzotrlazole, 2-(3'-tert-butyl-2'hydroxy-5'-(2-octyloxycarbonylethyl)phenyl)-benzotrlazole, 2-(3'-tert-butyl-5'-[2-(2-ethylhexyl-2-(3'-dodecyl-2'-hydroxy-5'-methyloxy)carbonylethyl]-2'-hydroxyphenyl)-benzotriazole, 2-(3'-tert-butyl-2'-hydroxy-5'-(2-isooctyloxycarbonylethyl)-phenylphenyl)-benzotriazole, 2,2'-methylenebls[4-(1,1,3,3-tetramethylbulyi)-6-benzotriazol-2-yi-phenol]; benzotriazole. transesterification product of 2-[3'-tert-butyl-5'-(2-methoxycarbonylethyl)-2'-hydroxyphenyl]benzotriazole with polyethylene glycol 300; $[R-CH_2CH_2-COO-CH_2CH_2]_{\infty}$ wherein R =

 $\label{eq:continuous} 3\mbox{-}4-\$

2.2. 2-Hydroxybenzophonones, for example the 4-hydroxy, 4-methoxy, 4-octyloxy, 4-decyloxy, 4-dedecyloxy, 4-benzyloxy, 4,2,4-irhydroxy or 2-hydroxy-4,4-dimethoxy derivative.

2.3. Esters of unsubstituted or substituted benzolo acids, for example 4-tert-butyl-phenyl sellcylate, phanyl saticytate, octylphenyl salicylate, dibenzoylresorcinol, bis(4-tort-butyl-benzoylresorcinol, benzoylresorcinol, 3,5-dl-tert-butyl-4-hydroxybenzolo acid 2,4-dl-tert-butyl-4-hydroxybenzolo acid hexadecyl ester, 3,5-dl-tert-butyl-4-hydroxybenzolo acid catacoyl ester, 3,5-dl-tert-butyl-4-hydroxybenzolo acid catacoyl ester, 3,5-dl-tert-butyl-4-hydroxybenzolo acid 2-mathw-4,8-dl-tert-butyl-4-hydroxybenzolo acid 2-mathw-4,8-dl-tert-butyl-4-hydroxybenz

2.4. Acrylates, for example α-cyano-β,β-diphanylacryllo solid ethyl ester or isooctyl ester, α-methoxycarbonylchnamio acid methyl ester, α-cyano-β-methyl-p-methoxychnamic acid methyl ester or butyl ester, α-methoxycarbonyl-β-methoxychnamic acid methyl ester, N-(β-methoxycarbonyl-β-cyanovinyl)-2-methyl-inddine.

2.5. Nickel compounds, for example nickel complexes of 2,2-4thio-bla[4-(1,1,3,3-birtz-methylbutyl)phenoi], such as the 1:1 or 1:2 complex, optionally with additional ligends, such as n-butylemine, trebtanotemine or N-oyalomov/dishanotemine, nickel dibutyl dittiocarbamate, nickel salts of 4-hydroxy-3,5-di-tert-butylbenzylphosphonic acid monosityl esters, such as of the methyl or ethyl seter, nickel complexes of ketoximes, such as of 2-hydroxy4-methylphenylundecyl ketoxime, nickel complexes of 1-phenyl4-leurcyl-5-hydroxyyarzois, optionally with additional ligands.

2.6. Stancally bindered amines, for example bis(2,2,6,6-tetramethylpiperid-4-yl) sebacate, bis(2,2,6,6-teiramethylpiperid-4-yl) succinate, bis(1,2,2,6,6-pentamethylpiperid-4-yl) sebacate, bls(1-octyloxy-2,2,6,6-tetramethylpipertd-4-yl) sebacate, n-butyl-3,5-dl-tert-butyl-4hydroxybenzylmalonic acid bis(1,2,2,6,6-pentamethylpiperidyl) ester, condensation product of 1-hydroxyethyl-2,2,6,6-tetramethyl-4-hydroxyplperidine and succinic acid, linear or cyclic condensation products of N,N'-bis(2,2,6,6-tetramethyl-4-piperidyl)hexamethylenedlamine and 4-tert-oct/lamino-2,6-dichloro-1,3,5-s-triazine, tris(2,2,6,6-tetramethyl-4-pipendyl) nitrilotriacetate, tetrakis(2,2,6,6-tetramethyl-4-piperidyl)-1,2,3,4-butanetetraoate, 1,1'-(1,2-ethanediyl)bls(3,3,5,5-tetramethylpiperazinone), 4-benzoyl-2,2,6,6-tetramethylpiperidine, 4-stearyloxy-2,2,6,6-tetramethylpiperidine, bis(1,2,2,6,6-pentamethylpiperidyl)-2-n-butyl-2-(2-hydroxy-3,5-di-tert-butylbenzyi) malonale, 3-n-octyl-7,7,9,9-tetramethyl-1,3,8-triazasplro[4.5]decane-2.4-dlone, bis(1-octyloxy-2,2,6,6-tetramethylpiperidyl) sebacate, bis(1-octyloxy-2,2,6,6-tetramethylpiperidyl) succinate, linear or cyclic condensation products of N,N'-bis(2,2,6,8-tetramethyl-4-piperidyl)hexamethylenediamine and 4-morpholino-2,6-dichloro-1,3,5-triazine, condensation product of 2-chloro-4,6-di(4-n-butylamino-2,2,6,6-tetramethylpiperidyi)-1,3,5-triazine and 1,2-bls(3-aminopropylamino)ethane, condensation product of 2-chloro-4,6-di(4-nbutylamino-1,2,2,6,6-pentamethylpiperidyl)-1,3,5-triazine and 1,2-bis(3-aminopropylamino)ethane, 8-acetyl-3-dodecyl-7,7,9,9-teiramethyl-1,3,8-triazaspiro[4.5]decane-2,4-dione, 3dodecyl-1-(2,2,6,6-tetramethyl-4-piperidyl)pyrrolidine-2,5-dione, 3-dodecyl-1-(1,2,2,6,6-pentamethyl-4-piperidyl)pyrrolldine-2,5-dione, mixture of 4-hexadecyloxy- and 4-steanyloxy2,2,6,6-tetramethylpiperidine, condensation product of N,N'-bls(2,2,6,6-tetramethyl-4-piperidyl)hexamethylenediamine and 4-cyclohexylamine-2,6-dichlore-1,3,5-triazine, condensation product of 1,2-bis(3-aminopropylamino)ethane and 2,4,6-trichloro-1,3,5-triazine and 4-butylamino-2,2,6,6-tetramethylpiperkline (CAS Reg. No. [136504-96-6]); condensation product of 1.6-diaminohexane and 2,4,6-trichloro-1,3,5-triazine and also N,N-dibutylamine and 4-butylamino-2,2,6,6-tetramethylpiperidine (CAS Reg. No. [192268-64-7]); N-(2,2,6,6-tetramethyl-N-(1,2,2,6,6-pentamethyl-4-piperidyl)-n-4-piperidyl)-n-dodecylsuccinimide, dodecylsuccinimide, 2-undecyl-7,7,9,9-tetramethyl-1-oxa-3,8-dlaza-4-oxo-spiro[4.5]decane, 7.7.9,9-tetramethyl-2-cycloundecyl-1-oxa-3,8-diaza-4of reaction product oxospiro[4.5]decane and epichiorohydrin, 1,1-bis(1,2,2,6,6-pentamethyl-4-piperidyl-N,N'-bis-formyl-N,N'-bis(2,2,6,6-tetramethyl-4oxycarbonyi)-2-(4-methoxyphenyl)ethene, piperidyl)hexamethylenediamine, diester of 4-methoxymethylenemalonic acid with 1,2,2,6,6poly[methylpropyl-3-oxy-4-(2,2,6,6-tetramethyl-4pentamethyl-4-hydroxypiperidine, piperidyi)]siloxane, reaction product of maleic anhydride o-olefin copolymer and 2,2,6,6tetramethyi-4-aminopiperidine or 1,2,2,6,6-pentamethyl-4-aminopiperidine.

2.7. Oxalio acid diamides, for example 4.4"-dioctyloxy oxanilide, 2,2"-diethoxy oxanilide, 2,2"-dioctyloxy-5,5"-di-tert-butyl oxanilide, 2,2"-dioceolyloxy-5,5"-di-tert-butyl oxanilide, 2-ethoxy-2"-ethyl oxanilide, N,N-big3-dimethylaminopropyl) oxalamide, 2-ethoxy-5-tert-butyl-2"-ethyl oxanilide and a mixture thereof with 2-ethoxy-2"-ethyl-5,4"-di-ter-butyl oxanilide, mixtures of o- and p-methoxy- and also of o- and p-ethoxy-di-substituted oxanilides.

2.8. 2-(2-thydroxynhenvt)-1.3.6-triazines, for example 2.4,6-tris(2-hydroxy-4-octyloxyphenyt)-1,3,5-triazine, 2-(2-thydroxy-4-octyloxyphenyt)-4,6-bis(2-thinethythenyt)-1,3,5-triazine, 2-(2,4-dihydroxyphenyt)-4,6-bis(2-thinethythenyt)-1,3,5-triazine, 2-(2-thydroxy-4-ctyloxyphenyt)-4,6-bis(2-thinethythenyt)-1,3,5-triazine, 2-(2-thydroxy-4-dotocyloxyphenyt)-4,6-bis(2-thinethythenyt)-1,3,5-triazine, 2-(2-thydroxy-4-tridocyloxyphenyt)-4,6-bis(2-thinethythenyt)-1,3,5-triazine, 2-(2-thydroxy-4-thydroxyphenyt)-4,6-bis(2-thinethythenyt)-1,3,5-triazine, 2-(2-thydroxy-4-(2-thydroxy-3-octyloxypropyloxyphenyt)-4,6-bis(2-thinethythenyt)-1,3,5-triazine, 2-(2-thydroxy-4-(2-thydroxy-3-octyloxypropyloxyphenyt)-4,6-bis(2-thinethythenyt)-1,3,5-triazine, 2-(4-thydroxy-4-(2-thydroxy-

(3-butoxy-2-hydroxypropoxy)phenyl]-1,3,5-triazine, 2-{2-hydroxyphenyl}-4-{4-mothoxyphenyl}-8-phenyl-1,3,5-triazino, 2-{2-hydroxy4-{3-{2-otliy(thexyl-1-oxy)-2-hydroxyy-1-y0xy)phenyl}-4,6-bis(2,4-dimethylphenyl)-1,3,5-triazine.

3. Metal deactivators, for exampte N,N-diphenyloxalic acid diamide, N-salicyloyl-hydrazine, N,N'-bia(salicyloyf)hydrazine, N,N'-bia(salicyloyf)hydrazine, N,N'-bia(salicyloyf)hydrazine, N,N'-bia(salicyloyf)hydrazine, 3-salicyloyfamino-1,24-trazole, bis(benzylidene)oxalic acid dihydrazide, oxaniide, isophthalic add dihydrazide, sobacic acid bis-phenylhydrazide, N,N'-diacofyladipic acid dihydrazide, N,N'-bis-salicyloyfoxalic acid dihydrazide, N,N'-bis-salicyloyfihopropionic acid dihydrazide.

4. Phosphites and phosphonites, e.g. triphenyl phosphile, diphenylalkyl phosphites, phenyldialkyl phosphiles, tris(nonylphenyl)phosphile, trilauryl phosphile, trioctadecyl phosphile, distearyl-pentaerythritoi diphosphite, tris(2,4-di-tert-butylphenyi)phosphite, diisodecylpeniaerythritol diphosphile, bis(2,4-di-tert-butylphenyl)pentaerythritol diphosphile, bis(2,4-dicumylphenyl)-pentaerythritol diphosphite, bis(2,6-di-tert-butyl-4-methylphenyl)pentaerythritol bis(2,4-di-tert-butyl-6bis-isodecyloxy-pentaerythritol diphosphite, diphosphite, diphosphile, bis(2,4,6-tri-tert-bulyiphenyl)pentaerythritol methylphenyl)pentaerythritol diphosphite, tristearyl sorbitol triphosphite, tetrakis(2,4-di-tert-butylphenyl)-4,4'-biphenylene diphosphonite, 6-isooctyloxy-2,4,8,10-tetra-tert-butyl-12H-dibenzo[d,g]-1,3,2-dioxaphosphooine, bis(2,4-di-tert-butyl-6-methylphenyi) methylphosphile, bis(2,4-di-tert-butyl-6-methylphenyl) ethylphosphile, 6-fluoro-2,4,8,10-tetra-tert-butyl-12-methyl-dibenzo[d,g]-1,3,2-dioxaphosphocine, 2,2',2"-nitrilo[triethyi-tris(3,3',5,5'-tetra-tert-butyl-1,1'-blphenyl-2,2'-diyl)-phosphite], 2-ethylhexyl-(3,3',5,5'-tetra-tert-butyl-1,1'-biphenyl-2,2'-diyl) phosphite, 5-butyl-5-ethyl-2-(2,4,6-tri-tert-bulylphenoxy)-1,3,2-dloxaphosphirane.

<u>6. Hvdroxylamines</u>, for example N,N-dibenzylitydroxylamine, N,N-diethythydroxylamine, N,N-dioctylhydroxylamine, N,N-dioctylhydroxylamine, N,N-dioctadecylitydroxylamine, N,N-dioctadecylhydroxylamine, N-hexadocyl-N-octadecyl-hydroxylamine, N-heptadocyl-N-octadecyl-hydroxylamine, N,N-dialkylitydroxylamine from hydroxylamines a tilliow fatty amines.

6. Niltones, for example N-benzyl-alpha-phenylnitrone, N-ethyl-alpha-methylnitrone, N-octyl-alpha-heptylnitrone, N-letradecyl-alpha-tridecylnitrone, N-tetradecyl-alpha-tridecylnitrone,

N-hexadecyl-alpha-pentadocylnitrone, N-ocladecyl-alpha-heptadecylnitrone, N-hexadecylalpha-heptadecylnitrone, N-ocladecyl-alpha-pentadecylnitrone, N-heptadecyl-alpha-heptadecylnitrone, N-ocladecyl-alpha-hexadecylnitrone, nifrones derived from N,N-dialkylhydroxylarninas prepared from hydrogenated taliow fatty amines.

- 7. Thiosyneralstic compounds, for example thiodipropionic acid dilauryl ester or thiodipropionic acid distearyl ester.
- 8. Peroxide-destroying compounds, for example esters of β-thio-dipropionic ecid, for example the leuryl, stearyl, myristly or trideoyl ester, merceptobenzimidazole, the zinc salt of 2-merceptobenzimidazole, zinc dibutyldithiocarbamate, dioctadeoyldisutifide, pentaerythritol tetraksigi declerolymerceptolypropionate.
- Polyamide stabilisers, for example copper salts in combination with lodides and/or phosphorus compounds and salts of divalent manganese.
- 10. Basio co-siabilisers, for example melamine, polyvinyfpyrrolidone, dicyandiamido, trially/ cyanurate, urea derivatives, hydrazine derivatives, amines, polyamides, polyurethanes, alkall metal and alkaline earth metal salts of higher fatty acids, for example calcium stearate, zino stearate, megnesium behenate, magnesium stearate, sodium ricinoleate, potassium palmitate, antimony pyrocatecholate or zino pyrocatecholate.
- 11. Nucleating agents, for example inorganic substances, e.g. talc, metal oxidea, such as titanium dioxide or magnesum oxide, phosphates, carbonates or suifates of preferably alkaline earth metals; organic compounds, such as mono-or poly-carboxylic acids and their salts, e.g. 4-tert-bulytibenzolc acid, adiple acid, diphanylacetic acid, sodium succinate or sodium benzoate; polymeric compounds, for example lonic copolymerisates ("lonomars"). Special preference is given to 1,3:2-4-bis(3/4-dimethytibenzylidene)sorbilol, 1,3:2-4-dif(paramethytidibenzylidene)sorbilol.
- 12. Fillers and reinforcing auents, for example calcium carbonate, silicates, glass fibres, glass beads, tatc, kaolin, mica, barlum sulfate, metal oxides and hydroxides, carbon black, graphilip, wood powders, and powders and fibres of other natural products, synthetic fibres.

13. Other additives, for example plasticisers, lubricants, emulsitiers, pigments, rheology additives, catalysts, flow improvers, optical brighteners, flame retardants, antistatics, blowing agents.

14. Benzofuranopos and Indolloness, for example as described in U.S. 4 325 863; U.S. 4 338 244; U.S. 5 175 312, U.S. 5 216 052; U.S. 5 252 643; DE-A-4 316 817; DE-A-4 316 876; EP-A-0 589 839 or EP-A-0 591 102, or 3-14-(2-sectoxyethoxy)phenyll-5-7-d-Ind-Indyl-5-errofuran-2-one, 5-7-d-Ind-Indyl-5-errofuran-2-one, 3-3-bis[57-d-Ind-Indyl-3-(4-[2-Inydroxyethoxy)phenyll-benzofuran-2-one, 3-4-lonexyphenyll-benzofuran-2-one, 3-4-d-Indxyyhenyll-5-7-d-Ind-Indyl-5-1-d-Indxyyhenyll-5-7-d-Indxy-benzofuran-2-one, 3-4-d-Indxyyhenyll-5-7-d-Indxy-benzofuran-2-one, 3-4-d-Indxy-benzofuran-2-one, 3-4-d-Indxy-benzofuran-2-one, 3-4-d-Indxy-benzofuran-2-one, 3-4-d-Indxy-benzofuran-2-one, 3-4-d-Indxy-benzofuran-2-one, 3-4-d-Indxy-benzofuran-2-one, 3-4-d-Indxy-benzofuran-2-one, 3-4-d-In

The costabilizers are added, for example, in concentrations of 0.01 to 10%, relative to the total weight of the synthetic polymer to be stabilized.

Preferred further additives are phenolic antioxidants, light-stabilizors, processing stabilizers, solvents, pigments, dyes, plasticizers, compatibilizers, toughening agents and/or thixotropic agents.

In addition to the nano fillers other fillers may be used as reinforcing agents (item 12 in the list), for example talc, calcium carbonate, hydrolatcite, mica, kaolin, motal hydroxides, especially aluminium hydroxide or magnesium hydroxide. These ere added to the synthetic polymers in concentrations, for example, of from 0.01 to 40 %, based on the overall weight of the synthetic polymers to be stabilized.

Carbon black as filler is added to the synthetic polymers in concentrations, judiclously, of from 0.01 to 5 %, based on the overall weight of the synthetic polymers to be stabilized.

Glass fibers as reinforcing agents are added to the synthetic polymers in concentrations, judiciously, of from 0.01 to 20 %, based on the overall weight of the synthetic polymers to be stabilized.

Further preferred compositions comprise in addition to components (a), (b) and (c) further additives as well, especially alkaline earth metal salts of higher fatty acids, for example calcium stearate.

As a conventional stabilizor combination for processing synthetic polymers, for example polyolellins, to form corresponding moukdings, the combination of a phenotic antioxidant with a secondary antioxidant based on an organic phosphile or phosphonlie is recommended.

Incorporation of components (b) and (c) and, if desired, further additives into the synthetic polymers is carried out by known methods, for example before or during moutding or else by applying the dissolved or dispersed compounds to the synthetic polymer, if appropriate with subsequent slow evaporation of the solvent.

The present invention also relates to a nanocomposite meterial in the form of a masterbalch or concentrate comprising component (a) in an amount of from 5 to 90%, component (b) in an amount of from 5 to 80%, and component (c) in an amount of from 0.5 to 50% by weight.

Components (b) and (c) and, if desired, further additives, can also be added before or during polymerisation or before crosslinking.

Components (b) and (o), with or without further additives, can be incorporated in pure form or encapsulated in waxes, oils or polymers into the synthetic polymer.

Components (b) and (c), with or without further additives, can also be sprayed onto the synthetic polymer. It is able to dilute other additives (for example the conventional additives indicated above) or their mells so that they too can be sprayed together with these additives onto the polymer. Addition by spraying on during the deactivation of the polymerization catalysis is perticularly advantageous, it being possible to carry out spraying using, for example, the steam used for deactivation.

In the case of spherically polymerized polyoletins it may, for example, be advantageous to apply components (b) and (c), with or without other additives, by spreying.

The synthetic polymers prepared in this way can be employed in a wide variety of forms, for example as foams, films, fibres, tepes, moulding compositions, as profiles or as binders for coating materials, especially powder coatings, adhesives, putties or especially as thick-layer polyclafin mouldings which are in long-term contact with extractive media, such as, for example, pipes for liquids or gases, films, fibres, geomembranes, tapes, profiles or tanks.

The preferred thick-layer polyolelin mouldings have a leyer thickness of from 1 to 50 mm, in particular from 1 to 30 mm, for example from 2 to 10 mm.

The compositions according to the invention can be advantageously used for the proparation of various shaped articles. Examples are:

- i-1) Floating devices, marine applications, pontoons, buoys, plastic lumber for decks, plans, boats, kayaks, oars, and beach reinforcements.
- 1-2) Automotive applications, in particular bumpers, deshboards, battery, rear and front linings, moldinge parts under the hood, hat shelf, trunk linings, interior linings, ale bag covers, electronic moldings for fittings (lights), panes for dashboards, headlamp glass, instrument panel, exterior linings, upholistery, automotive lights, head lights, parking lights, rear lights, stop lights, interior and exterior trims; door penels; gas tank; glazing front side; rear windows; seat backing, exterior penels, wire insulation, profile extrusion for sealing, eddiding, piller covers, chassis parts, exhaust systems, fuel filter / filler, fuel pumps, fuel tank, body side mouthings, convertible tops, exterior mirrors, exterior trim, fasteners / fixings, front end module, glass, hinges, lock systems, luggage / roof racks, prossed/stamped parts, seals, side impact protection, sound deadener? Insulator and sunroof.
- I-3) Road traffic devices, in particular sign postings, posts for road marking, car accessories, warning triangles, medical cases, helmets, tires.
- I-4) Devices for plane, rallway, motor car (car, motorbike) including furnishings.
- 1-5) Devices for space applications, in particular rockets and satellites, e.g. reentry shields.

- I-6) Devices for architecture and design, mining applications, acoustic quietized systems, street refuges, and shelters.
- II-1) Appliances, cases and coverings in general and electric-electronic devices (personal computer, telephone, portable phone, printer, television-sets, audio and video devices), flower pots, satellite TV bowl, and panel devices.
- II-2) Jacketing for other materials such as steel or textiles.
- II-3) Devices for the electronic industry, in particular insulation for plugs, especially computer plugs, cases for electric and electronic parts, printed boards, and materials for electronic data storage such as chips, check cards or credit cards.
- II-4) Electric appliances, in particular washing machines, tumblers, ovens (microwave oven), dish-washers, mixers, and frons.
- II-5) Covers for lights (e.g. street-lights, lamp-shades).
- II-6) Applications in wire and cable (semi-conductor, insulation and cable-jacketing).
- II-7) Foils for condensers, refrigerators, heating devices, air conditioners, encapsulating of electronics, semi-conductors, coffee machines, and vacuum cleaners.
- iii-1) Technical articles such as cogwheel (gear), slide fittings, spacers, screws, bolts, handles, and knobs.
- III-2) Rotor blades, ventilators and windmill vanes, solar devices, swimming pools, swimming pool covers, pool liners, pond liners, closets, wardrobes, dividing walls, stat walls, folding walls, roofs, shutters (e.g. roller shutters), fittings, connections between pipes, sleaves, and conveyor betts.
- III-3) Sanitary articles, in particular shower cubicles, lavatory seats, covers, and sinks.

- III-4) Hygienic articles, in particular diapers (bables, adult incontinence), feminine hygiene articles, shower curtains, brushes, mats, tubs, mobile toilets, tooth brushes, and bed pans.
- III-5) Pipes (cross-linked or not) for water, waste water and chemicals, pipes for wire and cable protection, pipes for gas, oil and sewage, guttering, down pipes, and drainage systems.
- III-6) Profiles of any geometry (window panes) and siding.
- III-7) Glass substitutes, in particular extruded plates, glazing for buildings (monolithic, twin or multiwall), aircraft, schools, extruded sheets, window film for architectural glazing, train, transportation, sanitary articles, and greenhouse.
- III-8) Plates (walls, cutting board), extrusion-coating (photographic paper, tetrapack and pipe coating), silos, wood substitute, plastio lumber, wood composites, walls, surfaces, furniture, decorative foil, floor coverings (interior and exterior applications), flooring, duck boards, and titles.
- III-9) Intake and outlet manifolds.
- III-10) Cement-, concrete-, composite-applications and covers, siding and cladding, hand ralls, banisters, kitchen work tops, roofing, roofing sheets, tiles, and tarpaulins.
- IV-1) Plates (walls and cutting board), trays, artificial grass, astroturf, artificial covering for stadium rings (athletics), artificial floor for stadium rings (athletics), and tapes.
- IV-2) Woven fabrics continuous and staple, fibers (carpots / hygienic articles / geotextiles / monofilaments; fillers; wipes / curtains (shades) / medical applications), bulk fibers (applications such as gown / protection olothes), nets, ropes, cables, strings, cords, threads, safety seat-belts, clothes, underwear, gloves; boots; nuber boots, Intimate apparel, garments, swimwear, sportswear, umbrellas (parasol, sunshado), parachutes, paraglides, salts, 'balloon-slik', camping articles, lents, aifbeds, sun beds, bulk bags, and bags.

IV-3) Membranes, Insulation, covers and seals for roofs, tunnels, dumps, ponds, dumps, walls roofing membranes, geomembranes, extraming pools, curtains (shedes) / sun-shields, awarlings, canoples, wallpaper, food packing and wrapping (flexible and solid), medical packeging (flexible & solid), airbagslanfely belts, erm- and head rests, carpets, centre console, dashboard, cockpits, door, overhead console module, door trim, headliners, interior lighting, interior mirrors, percel shelf, rear luggage cover, seats; steering column, steering wheel. facilies and trunk trim.

V) Films (packaging, dump, laminating, agriculture and horticulture, greenhouse, mulch, tunnel, silage), bale wrap, swimming pools, waste bags, wallpaper, stretch film, raffla, desalhation film, batteries, and connectors.

VI-1) Food packing and wrapping (flexible and solid), bottles.

VI-2) Storage systems such as boxes (crates), luggage, chest, household boxes, pallets, shelves, tracks, screw boxes, packs, and cans.

VI-3) Cartridges, syringes, medical applications, containers for any transportation, waste baskets and waste bins, waste bags, bins, dust bins, bin liters, wheely bins, container in general, tanks for water / used water / chemistry / gas / oil / gasoline / disset; tank liners, boxes, crates, battery cases, troughs, medical devices such as piston, ophthalmic applications, diagnostic devices, and packing for pharmacouticals bitster.

VII-1) Extrusion coating (photo paper, tetrapack, pipe coating), household articles of any kind (e.g. appliances, thermos bottle I clothes hanger), fastoning systems such as plugs, wire and cable clamps, zippers, closures, locks, and snap-closures.

VII-2) Support devices, articles for the leisure time such as sports and filness devices, gymnastics mals, ski-boots, inline-skales, skis, big foot, athletic surfaces (e.g. tennis grounds); screw tops, tops and stoppers for bottles, and cans.

VII-3) Furniture in general, foamed articles (cushions, impact absorbers), foams, sponges, dish clothes, mats, garden chairs, stadium seats, tables, couches, toys, building kits (boards / figures / balls), playhouses, sildes, and play vehicles. VII-4) Materials for optical and magnetic data storage.

VII-5) Kitchen ware (eating, drinking, cooking, storing).

VII-6) Boxes for CD's, cassettes and video tapes; DVD electronic articles, office supplies of any kind (ball-point pens, stamps and ink-pads, mouse, shelves, tracks), bottles of any volume and content (drinks, detergente, cosmetics including perfumes), and adhesive tapes.

VII-7) Footwear (shoes / shoe-soles), insoles, spats, adheelves, structural adheelves, food boxes (fruit, vegetables, meat, fish), synthetic paper, tabels for bottles, couches, artificial joints (human), printing plates (flexographic), printed circuit boards, and display technolooles.

VII-8) Devices of filled polymers (talc, chalk, china olay (kaolin), wollastonite, pigments, carbon black, TiO₂, mica, nanocomposites, dolomite, silicates, glass, asbestos).

Thus, a further embodiment of the present invention relates to a shaped article, in particular a film, pipe, profile, bottle, tank or container, filter containing a composition as described above.

A further embodiment of the present invention relates to a molded article containing a composition as described above. The molding is in particular effected by injection, blow, compression, roto-molding or elush-molding or extrusion.

The present invention also relates to a process for the preparation of a synthetic polymer nanocomposite material which comprises melt mixing a mixture of a) a synthetic polymer, b) a filler, and o) a dispersing agent prepared by controlled free radical polymerization (CFRP).

The melt mixing can be carried out in any heatable container equipped with a slirrer, for example in a closed apparatus such as a knoader, mixer or stirred vessel. The incorporation is preferably carried out in an extruder or in a knoader. It is immatical whether processing takes place in an inert atmosphere or in the presence of oxygen. The addition of components (a), (b) and (c) can be carried out in all customary mixing machines in which the polymer is melted and mixed with the additives. Suitable machines are known to those skilled in the art. They are predominantly mixers, kneaders and extruders. The process is preferably carried out in an extruder by introducing the additive during processing. Particularly preferred processing machines are single-screw extruders, contrarotaling and corotating twin-screw extruders, planetary-gear extruders, ring extruders or cokneaders, it is also possible to use processing machines provided with at least one gas removal compartment to which a vacuum can be applied. Suitable extruders and kneaders are described, for example, in Handbuch der Kunststoffextrusion, Vol. 1, Grundlagen, Editors F. Hensen, W. Knappe, H. Potente, 1989, pp. 3-7, ISBN:3-446-14339-4; and Vol. 2 Extrusionsanlagen 1986, ISBN 3-446-14329-7. For example, the screw length is 1 - 60 screw diameters, preferably 35 - 48 screw diameters. The rotational speed of the screw is preferably 10 to 600 rotations per minute (rpm), for example 25 - 300 rpm. The maximum throughput is dependent on the screw diameter, the rotational speed and the driving force. The process of the present invention can also be carried out at a level lower than meximum throughput by varying the parameters mentioned or employing weighing machines delivering dosage amounts, if a plurality of components are added, these can be premixed or added individually.

Also of Interest is a process for the preparation of a synthetic polymer nanocomposite material, wherein the melt mixing of the components (synthetic polymer, filler and dispersing agent prepared by controlled free radical polymertzation) occurs between 120 and 290°C, preferably between 140 and 250°C, for example between 170 and 230°C.

The present invention also relates to synthetic polymer nanocomposites obtained by the above mentioned process.

The preferred components (b) and (c), and optionally futher additives, in the process for the preparation of a synthetic polymer nanocomposite material are the same as those described for the composition.

A preferred embodiment of the present invention is also the use of a dispersing agent prepared by controlled free radical polymerization to intercelate and exfoliete a filter and disperse the filter in a synthetic polymer matrix to form a nanocomposite material. The preferred dispersing agent, filler and synthetic polymer, and optionally further additives, for this use are the same as those described for the composition.

The following examples illustrate the invention further. Parts or percentages relate to weight.

Example 1: Preparation of dispersing agents by controlled free radical polymerization,

The preparation and characterization of block or comb copolymers by controlled free radical polymerization is disclosed for example in C. Auschra et al., Progress in Organic Coeffings 45, 83 — 93 (2002). Based on these procedures the following block copolymers are prepared with the properties disclosed in Table 1.

| Table 1: | | | | | | |
|----------------------------|--------------------|---------------------------------|----------------------|---------------------------------|------------------------------|--|
| Dispersing agent (Example) | Block 1 | Block 1 length ¹⁾ | Block 2 | Block 2 length ²⁾ | M _n ³⁾ | M _e /M _e ⁴⁾ |
| 1a | PBA ⁶⁾ | 58 | PDMAEA ⁶⁾ | 16 | 7400 | 1.44 |
| 1b | PBA ⁵⁾ | 100 | PDMAEA ⁶⁾ | 30 | 11200 | 1.55 |
| 1c | PBMA ⁷⁾ | 66 | PDMAEA ⁶⁾ | 25 | n.d. ⁸⁾ | n.d. ⁸⁾ |

- 1) Average number of monomer units of the first block.
- Average number of monomer units of the second block.
- 3) Molecular weight obtained from ¹HNMR integration.
- 4) Polydispersity of polymer obtained from GPC with THF as solvent.
- PBA is poly(butyl acrylate).
- 6) PDMAEA is poly(dimethylaminoethyl acrylate).
- 7) PBMA is poly(butyl methacrylate).
- 8) n.d. means "not determined".

Example 2: Preparation of polypropylene nanocomposites in a batch mixer based on unmodified sodium montmorillonite.

50 g of polypropylene [Basell KY 6100 (RTM)], is blended with 0.25 % of Irganox 1010 (RTM) (ponteerythriot) etrakeig-(3,5-dl-4rr-butyl-4-hydroxyphenyf)propionate)) and 0.25 % of Irgafos 168 (RTM) (trieg-2,4-ther-butyl-henyl-phosphite), 5 % of a monthrorillonite clay [Cicloite (Na*) (RTM) obtained from Southern Clay Industries] and 2.5 % of a dispersing agent prepared by controlled free radical polymerisation (CFRP) according to Table 1 in a plastic cup and then added to a batch mixer operating at 25 rpm and 180°C, the mixture is then taken to 50 rpm for 10 minutes. Small seale injection moutding is performed with a CS-168 MMX Minimax moulder. The mixing chamber is preheated to 230°C and the mould is oven heated to 120°C for one hour prior to injection moutding. Approximately 5 g of the material is placed in the mixing chamber and heated for 4 to 5 minutes. A small shot is taken form the CS-183 MMX Minimax moulder prior to placing the mould in position. Six tensils bars with the dimensions 18 mm by 5 mm by 0.85 mm are produced for each of the 5 ametrial.

Tensile testing is performed according to ASTM D 638 with a Rheometrics mini material tenelle tester [Minimat 2000 (RTM)] equipped with a 1000 N load cell (40 mm/min). The results are summarized in Table 2.

Table 2

| 18018 2: | | | | | |
|------------------|----------------------|--------------------------|--|--|--|
| Example | Dispersing agent | d-001 in A ^{c)} | Normalized Modulus of Elasticity ⁶⁾ | Elongation at break (%) ^{e)} | |
| 2a ⁶⁾ | none (and no filler) | _ | 1.00 | 1.00 | |
| 2b*) | None | 10 | 1.12 | 0.97 | |
| 20 ^{b)} | Example 1a | 13 | 1.32 | 1.30 | |
| 2d ^{b)} | Example 1b | 13 | 1.04 | 1.26 | |
| 2e ^{b)} | Example 1c | n.d. ⁰ | n.d. ⁰ | n.d. ⁰ | |

- a) Comparative Example.
- b) Example according to the invention.
- c) Interlayer distance determined by X-ray diffraction.
- Normalized modulus of elasticity relative to polypropylene processed under similar conditions (= 1.0).
- Normalized elongation at break relative to polypropylene processed under similar conditions (= 1.0).
- f) n.d. means "not determined".

The X-ray diffraction (XRD) spectra show that all copolymers synthesized via CFRP eccording to Table 2 intercalate the used diay in the direct malt blending experiments in the batch maker. This is demonstrated by an increase in d-spacing of from 10 Å (comparative Example 2b) to 13 Å (Examples 2c and 2d according to the invention). The elasticity modulus of samples is anhanced by 4-32% with respect to polypropylene without dispersing agent and/or fillers. Surprisingly, the ofengation at break is also enhanced by 26-30%.

Example 3: Preparation of acrylic block or gradient copolymers by RAFT polymerization.

a) Preparation of poly(butyl acrylate) (PBA).

A solution 200 g (1.56 mol) of bulyl acrylete, 21.3 g (0.056 mol) of S-dodecyl S'-phenylethyl frithicarbonate) and 915 mg (0.0056 mol) of AIBN in 200 ml of dry toluene is depassed by purging with ergon for 3 hours. Polymerization is carried out at 70 °C with stifring under argon for 20 hours (*1 h MMr: conversion 55%; GPC: Mn 3700, Mw/Mn 1.16).

b) Preparation of poly(butyl acrylate)-block-poly(PEGMEA) (PBA-b- PPEGMEA).

A solution of 78.7 g (0.173 mol) of methyl (no)yethylene glycol) monoacrylate (average DP=3.1) and 915 mg (0.0056 mol) of AIBN in 100 ml of dry toluene is degassed by purging with argon for 3 hours. This solution is added to the above solution [Example 39] by syringe. The mixture is stirred at 60°C under argon for 22 hours. The solvent is removed by evaporation under vacuum. (Viscous yellow oit; "H NMR: total conversion ~100%; GPC: Mn 4400, MwWin 1.15).

c) Preparation of poly(buty) acrylate)-block-poly(DEGEEA) (PBA-b- PDEGEEA).

A solution of 39.2 g (0.208 mol) of disinlyience glycol) eithy either acrylate and 916 mg (0.0058 mol) of AlBN in 100 ml of dry toluone is degessed by purging with ergon for 3 hours. This solution was added to the above solution [Example 3a] by syrings. The mixture is stirred at 60°C under argon for 48 hours. The solvent is removed by evaporation under vacuum. (Viscous yellow out; "In NMR: total conversion ~100%; GPC: not available due to incomplete solubility in THF).

d) Preparation of poly(MMA-grad- PEGMEA).

A solution of 24.0.0 g (2.392 mol) of motityl methacrylate, 6.0.0 g (0.132 mol) poly(ellrylane glycol) methyl ether acrylate, 22.9 g (0.060 mol) of S-dodecyl S'-phenylathyl trititiocarbonate) and 1.47 g (0.0090 mol) of AIBN in 300 ml of dry fotuene is degassed by purging with argon for 3 houre. The mixture is ethred at 70°C under argon for 48 hours. The polymer is collected by precipitation into methanol. (yellow solid; "H NMR; total convension ~100%; GPC: Mn 7100. MwMRh 1.47).

e) Preparation of poly(MMA-grad- DEGEEA).

A solution of 24.0.0 g (2.392 mol) of methyl mothacrylate, 60.0 g (0.319 mol) dijethylene glycol) ethyl athor acrylate, 22.9 g (0.000 mol) of S-dodecyl S-phanylethyl trithiccarbonate) and 885 mg (0.0000 mol) of AIBN in 300 ml of dry toluene is degassed by purging with argon for 3 hours. The mbdure is stirred at 70°C under argon for 48 hours. The polymer is collected by precipitation into methanol. (viscous yellow oil; "H NIMR: total conversion ~99%; GPC; Mn 8100, MwMn 1.50).

Table 3

| Tubia or | | | | | | |
|----------------------------|--------------------------------|--------------------------------|-----------------------|---------------------------------|------------------------------|--|
| Dispersing agent (Example) | Block 1 | Block 1 length ⁹ | Block 2 | Block 2 length ²⁾ | M _n ³⁾ | M _a /M _n ⁴⁾ |
| 3b | PBA ⁶⁾ | 25.2 | PPEGMEA ⁶⁾ | 2.8 | 4400 | 1.15 |
| 30 | PBA ⁶⁾ | 28 | PDEGEEA ⁷⁾ | 3.1 | 4600 | - |
| 3d | PMMA-grad-PEGMEA ⁸⁾ | 56.6 | - | | 7100 | 1.47 |
| 3e | PMMA-grad-DEGEEA8) | 42.5 | - | | 5100 | 1.50 |

- 1) Average number of monomer units of the first block.
- 2) Average number of monomer units of the second block.
- Molecular weight obtained from ¹HNMR integration.
- 4) Polydispersity of polymer obtained from GPC with THF as solvent.
- 5) PBA is poly(butyl acrylate).
- PPEGMEA is poly(poly(ethylene glycol) methyl acrylate).
- 7) PDEGEEA is poly(diethylene glycol ethyl acrylate).
- PMMA-grad-DEGEEA is poly(methyl methacrylate-grad-diethylene glycol ethyl acrylate) gradlent copolymer (weight ratio MMA:DEGEEA 4:1)
- PMMA-grad-PEGMEA is poly(methyl methacrylate-grad-[poly(othylene glycol) methyl acrylate)]) gradient copolymer (weight ratio MMA: PEGMEA 4:1)

Example 4: Preparation of polypropylene nanocomposites in a twin screw extruder.

Processing is carried out with a Japan Steel Works 30 mm diameter twin screw extruder of L/D ratio 42 (JSW TEX 30) that comprises ten temperature controlled barrel sections each

with L/D of 3.5, three unheated sampling zones with L/D 1.167, and a cooled feed block with L/D 3.5. The screw configuration consists of a combination of mixing, kneeding almonveying elements familiar to bose skilled in the art. Materials are fed into the extruder via a JSW TTF20 gravimetric feeder (Feed 1) and a K-Tron KOX gravimetric feeder (Feed 2). The JSW TEX 30 is operated in a co rotating (Intermeshing self wiphraj) mode with throughput of 10 kg/hr and a screw speed of 200 rpm. Vacuum ventling is applied to the final barrel section. The extrudato is cooled in a water filled strand bath and pelletized.

in a first stop a 10 wt% day masterbatch is prepared. Feed 1 comprises a dry blend of polypropylene [Basel HF400N (RTM)] and a stabilizer which is composed of 0.25 wt% of inganox 1010 (RTM) (pentaerythritol letraktis[3-43,6-41-ert-butyl-4-hydroxyphenyl)propoloratel) and 0.25 wt% of Ingalos 168 (RTM) (intsi2.4-di-tert-butylphenyl)phosphita). Feed 2 comprises a blend of the day [Cloielle (Na*) (RTM) obtained from Southern Clay Industries] and the non-ionic surfactant in the ratio defined in Table 4. All barrel sections are heated to 170°C.

In the second step the masterbatch is let down to the required clay level (see Table 4) by blending the masterbatch (Feed 2) with further polypropylene plus stabilizer (Feed 1). The first barrel section is heated at 180°C the remaining barrel sections are heated at 200°C.

Injection moulding of the extruded samples are preformed with a Gindonati Milacron VS55 28 mm diameter Injection moulding machine comprising four temperature controlled sections of L/D23/1. The machine is operated at a clamp force of 50 tens and at a maximum injection pressure of 2005 bar.

Tensile testing is performed according to ISO 521 using an instron 5500R material tensile tester. Tensile properties (vs polypropylene) for nanocomposite prepared with a ratio of copolymer; clay of 1:5 are summarized in Tables 4 and 5.

| Ta | | |
|----|--|--|
| | | |
| | | |

| Example | Amount of Cloisite | Copolymer | Tensile Strength (MPa)) | Modulus of Elasticity (MPa) ⁰⁾ | Elongation at Break (%) |
|------------------|-----------------------|---------------|-------------------------------|---|----------------------------|
| 4a°) | none | none | 26.3 | 1470 | >800 |
| 4b ^{b)} | 5% | 1% Example 3b | 26.9 | 1560 | 106 |
| 40 ^{b)} | 5% | 1% Example 3c | 26,8 | 1540 | 141 |
| 4d ^{b)} | 5% | 1% Example 3d | 26.3 | 1530 | . 34 |
| 4e ^{b)} | 5% | 1% Example 3e | 26.5 | 1590 | 31 |
| 4f ^{b)} | 5% | 1% Example 1a | 26.8 | 1520 | 180 |
| 4g ^{b)} | 5% | 1% Example 1c | 26.6 | 1510 | 140 |

Table 5:

| | Example | Amount of Cloisite | Copolymer | Tensile Strength (MPa)) | Modulus of Elasticity (MPa) ^{d)} | Elongation at Break (%) |
|---|------------------|-----------------------|---------------|-------------------------------|---|----------------------------|
| ı | 5a*) | none | none (| 25.5 | 1470 | >800 |
| 1 | 5b ^{b)} | 1% | 1% Example 1a | 25.5 | 1826 | 49 |

- a) Comparative Example.
- b) Example according to the invention.

All examples of Table 4 demonstrate that the mechanical properties can be improved compared to polypropylene by using copolymens prepared by controlled radical polymerization according to the present invention.

What is claimed is:

- 1. A composition comprising
 - a) a synthetic polymer,
 - b) a filler, and
 - c) a dispersing agent prepared by controlled free radical polymerization (CFRP).
- 2. A composition according to claim 1, wherein component (a) is a polyolefin.
- 3. A composition according to claim 1, wherein component (b) is a filler in nanoparticles.
- A composition according to claim 1, wherein component (b) is a natural or synthetic
 phyllosilicate or a mixture of such phyllosilicates.
- 5. A composition according to claim 1, wherein component (b) is a layered silicate clay.
- 6. A composition according to claim 1, wherein component (b) is a montmorillonite, bentonite, beldelite, mica, hectorite, saponite, nontronite, sauconite, vermiculite, ledikite, magadite, kenyalfe, stevensite, volkonskotte, hydrotalcite or a mixture thereof.
- A composition according to claim 1, wherein component (c) is a random, block, star or comb copolymer prepared by controlled free radical polymerization.
- 8. A composition according to claim 1, wherein component (c) is a block or comb acryllo copolymer prepared by controlled free radical polymerization.
- 9. A composition according to claim 1, wherein component (b) is present in an amount of from 0.1 to 40 %, based on the weight of component (a).
- 10. A composition according to claim 1, wherein component (c) is present in an amount of from 0.1 to 20 %, based on the weight of component (a)

- A composition according to claim 1, comprising in addition, besides components (a), (b) and (c), further additives.
- 12. A composition according to claim 11, comprising as further additives phenolic anticod-dants, light-stabilizers, processing stabilizers, solvents, pigments, dyes, plasticizers, compatibilizers, toughening agents and/or thirotropic agents.
- 13. A composition according to claim 1 in the form of a masterbatch or concentrate comprising component (a) in an amount of from 5 to 80 %, component (b) in an amount of from 5 to 80 %, and component (c) in an amount of from 0.5 to 50 % by weight.
- 14. A process for the preparation of a synthetic polymer nanocomposite material which comprises melt mixing a mixture of a) a synthetic polymer, b) a filler, and o) a dispersing seent prepared by controlled free radical polymer/sation (CFRP).
- 15. A process according to claim 14, wherein the melt mixing occurs between 120 and $290^{\circ}\mathrm{C}$.
- 16. A synthetic polymer nanocomposite obtained by a process according to claim 14.
- 17. Use of a dispersing agent prepared by controlled free radical polymerization to intercalate and exfoliate a filter and disperse the filter in a synthetic polymer matrix to form a nano-composite material.

INTERNATIONAL SEARCH REPORT



Relevant to claim No.

A, CLASSIFICATION OF SUBJECT MATTER IPC 7 C07K14/665 C07K1/00

C. DOCUMENTS CONSIDERED TO BE RELEVANT

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum decumentation searched (classification system followed by classification symbols)

Category * Citation of document, with indication, where appropriate, of the relevant passages

Documentation searched other than minimum documentation to the extent that such documents are included. In the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data, PAJ, MEDLINE, BIOSIS, EMBASE, CHEM ABS Data

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| | NL - 2200 HV HSWIK Tel. (+31-70) 340-2040, Tx. 31 651 apo nl, Fax: (+31-70) 340-3016 | Schmidt, H | |

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